#### HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use SELZENTRY safely and effectively. See full prescribing information for SELZENTRY.

SELZENTRY (maraviroc) tablets, for oral use SELZENTRY (maraviroc) oral solution Initial U.S. Approval: 2007

#### WARNING: HEPATOTOXICITY

See full prescribing information for complete boxed warning.

- Hepatotoxicity has been reported which may be preceded by severe rash or other features of a systemic allergic reaction (e.g., fever, eosinophilia, or elevated IgE). (5.1)
- Immediately evaluate patients with signs or symptoms of hepatitis or allergic reaction. (5.1)

#### ------ INDICATIONS AND USAGE-----

SELZENTRY is a CCR5 co-receptor antagonist indicated in combination with other antiretroviral agents for the treatment of only CCR5-tropic HIV-1 infection in patients 2 years of age and older weighing at least 10 kg. (1) Limitations of Use:

Not recommended in patients with dual/mixed- or CXCR4-tropic HIV-1.

#### -----DOSAGE AND ADMINISTRATION -----

Prior to initiation of SELZENTRY, test all patients for CCR5 tropism using a highly sensitive tropism assay. (2.1)

SELZENTRY tablets and oral solution are taken twice daily by mouth and may be taken with or without food. SELZENTRY must be given in combination with other antiretroviral medications. (2.2)

Recommended Dosage in Adults: (2.3)

	Dosage of
Concomitant Medications	SELZENTRY
When given with potent CYP3A inhibitors (with or	150 mg
without potent CYP3A inducers) including PIs	twice daily
(except tipranavir/ritonavir), delavirdine (2.3, 7.1)	
With NRTIs, tipranavir/ritonavir, nevirapine,	300 mg
raltegravir, and other drugs that are not potent	twice daily
CYP3A inhibitors or CYP3A inducers (2.3, 7.1)	
With potent CYP3A inducers including efavirenz	600 mg
(without a potent CYP3A inhibitor) (2.3, 7.1)	twice daily

A more complete list of coadministered drugs is listed in *Dosage and Administration*. (2)

Pediatric Patients Aged 2 Years and Older and Weighing at Least 10 kg: Administer twice daily. Dosage should be based on body weight (kg) and concomitant medications and should not exceed the recommended adult dose.

Patients with Renal Impairment: Dose adjustment may be necessary in patients with renal impairment. (2.5)

#### ----- DOSAGE FORMS AND STRENGTHS-----

- Tablets: 25 mg, 75 mg, 150 mg and 300 mg. (3)
- Oral Solution: 20 mg per mL (3)

#### ----- CONTRAINDICATIONS -----

 SELZENTRY is contraindicated in patients with severe renal impairment or end-stage renal disease (ESRD) (CrCl less than 30 mL per minute) who are concomitantly taking potent CYP3A inhibitors or inducers. (4)

#### ----- WARNINGS AND PRECAUTIONS -----

- Hepatotoxicity accompanied by severe rash or systemic allergic reaction, including potentially life-threatening events, has been reported. Hepatic laboratory parameters including ALT, AST, and bilirubin should be obtained prior to starting SELZENTRY and at other time points during treatment as clinically indicated. If rash or symptoms or signs of hepatitis or allergic reaction develop, hepatic laboratory parameters should be monitored and discontinuation of treatment should be considered. When administering SELZENTRY to patients with pre-existing liver dysfunction or who are co-infected with hepatitis B and/or C virus, additional monitoring may be warranted. (5.1)
- Severe and potentially life-threatening skin and hypersensitivity reactions
  have been reported in patients taking SELZENTRY. This includes cases
  of Stevens-Johnson syndrome, hypersensitivity reaction, and toxic
  epidermal necrolysis. Immediately discontinue SELZENTRY and other
  suspected agents if signs or symptoms of severe skin or hypersensitivity
  reactions develop and monitor clinical status, including liver
  aminotransferases, closely. (5.2)
- More cardiovascular events, including myocardial ischemia and/or infarction, were observed in treatment-experienced subjects who received SELZENTRY. Additional monitoring may be warranted. (5.3)
- If patients with severe renal impairment or ESRD receiving SELZENTRY (without concomitant CYP3A inducers or inhibitors) experience postural hypotension, the dose of SELZENTRY should be reduced from 300 mg twice daily to 150 mg twice daily. (5.3)

#### ----- ADVERSE REACTIONS -----

- The most common adverse events in treatment-experienced adult subjects (greater than 8% incidence) which occurred at a higher frequency compared with placebo are upper respiratory tract infections, cough, pyrexia, rash, and dizziness. (6.1)
- The most common adverse events in treatment-naive adult subjects (greater than 8% incidence) which occurred at a higher frequency than the comparator arm are upper respiratory tract infections, bronchitis, flatulence, bloating and distention, upper respiratory tract signs and symptoms, and gastrointestinal atonic and hypomotility disorders. (6.1)
- The most common adverse reactions in treatment-experienced pediatric subjects (greater than or equal to 3% incidence) are vomiting, abdominal pain, diarrhea, nausea, and dizziness. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact ViiV Healthcare at 1-877-844-8872 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

# ----- DRUG INTERACTIONS-----

- Coadministration with CYP3A inhibitors, including protease inhibitors (except tipranavir/ritonavir) and delavirdine, will increase the concentration of SELZENTRY. (7.1)
- Coadministration with CYP3A inducers, including efavirenz, may decrease the concentration of SELZENTRY. (7.1)
- Coadministration with St. John's wort is not recommended. (7.1)

#### ----- USE IN SPECIFIC POPULATIONS -----

• Lactation: Women infected with HIV should be instructed not to breastfeed due to the potential for HIV transmission. (8.2)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.

Revised: 07/2018

#### **FULL PRESCRIBING INFORMATION: CONTENTS\***

#### WARNING: HEPATOTOXICITY

INDICATIONS AND USAGE

#### 2 DOSAGE AND ADMINISTRATION

- Testing prior to Initiation of SELZENTRY
- General Dosing Recommendations
- Recommended Dosage in Adults
- Recommended Dosage in Pediatric Patients 2.4
- Recommended Dosage in Patients with Renal Impairment

#### DOSAGE FORMS AND STRENGTHS

CONTRAINDICATIONS

#### WARNINGS AND PRECAUTIONS

- Hepatotoxicity
- Severe Skin and Hypersensitivity Reactions 5.2
- 5.3 Cardiovascular Events
- Immune Reconstitution Syndrome 5.4
- Potential Risk of Infection 5.5
- Potential Risk of Malignancy 5.6

#### ADVERSE REACTIONS

- Clinical Trials Experience 6.1
- Postmarketing Experience

#### DRUG INTERACTIONS

Effect of Concomitant Drugs on the Pharmacokinetics of Maraviroc

#### USE IN SPECIFIC POPULATIONS

- Pregnancy
- 8.2 Lactation
- Pediatric Use 8.4
- 8.5 Geriatric Use
- 8.6 Renal Impairment
- 8.7 Hepatic Impairment

#### 10 OVERDOSAGE

#### 11 DESCRIPTION

#### 12 CLINICAL PHARMACOLOGY

- 12.1 Mechanism of Action
- 12.2 Pharmacodynamics
- 12.3 Pharmacokinetics

# 12.4 Microbiology 13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
14 CLINICAL STUDIES

- 14.1 Clinical Studies in Adult Subjects14.2 Clinical Studies in Pediatric Subjects
- 15 REFERENCES
- 16 HOW SUPPLIED/STORAGE AND HANDLING
- 17 PATIENT COUNSELING INFORMATION

\*Sections or subsections omitted from the full prescribing information are not listed.

#### FULL PRESCRIBING INFORMATION

2	WARNING: HEPATOTOXICITY
3	Hepatotoxicity has been reported with use of SELZENTRY. Severe rash or evidence of a
4	systemic allergic reaction (e.g., fever, eosinophilia, or elevated IgE) prior to the
5	development of hepatotoxicity may occur. Patients with signs or symptoms of hepatitis or
6	allergic reaction following use of SELZENTRY should be evaluated immediately [see
7	Warnings and Precautions (5.1)].

#### 8 1 INDICATIONS AND USAGE

- 9 SELZENTRY is indicated in combination with other antiretroviral agents for the treatment of
- only CCR5-tropic human immunodeficiency virus type 1 (HIV-1) infection in patients 2 years of
- age and older weighing at least 10 kg.
- 12 Limitations of Use:

1

• SELZENTRY is not recommended in patients with dual/mixed- or CXCR4-tropic HIV-1 [see Microbiology (12.4)].

#### 15 2 DOSAGE AND ADMINISTRATION

#### 16 **2.1** Testing prior to Initiation of SELZENTRY

- 17 Prior to initiation of SELZENTRY, test all patients for CCR5 tropism using a highly sensitive
- tropism assay. SELZENTRY is recommended for patients with only CCR5-tropic HIV-1
- 19 infection. Outgrowth of pre-existing low-level CXCR4- or dual/mixed-tropic HIV-1 not detected
- 20 by tropism testing at screening has been associated with virologic failure on SELZENTRY [see
- 21 *Microbiology* (12.4), *Clinical Studies* (14.1)].
- 22 Monitor patients for ALT, AST, and bilirubin prior to initiation of SELZENTRY and at other
- 23 time points during treatment as clinically indicated [see Warnings and Precautions (5.1)].

# 24 **2.2** General Dosing Recommendations

- SELZENTRY tablets and oral solution are taken twice daily by mouth and may be taken with or without food.
- SELZENTRY must be given in combination with other antiretroviral medications.
- The recommended dosage of SELZENTRY differs based on concomitant medications due to drug interactions.

# 30 2.3 Recommended Dosage in Adults

- 31 Table 1 displays oral dosage of SELZENTRY based on different concomitant medications [see
- 32 Drug Interactions (7.1)].

#### **Table 1. Recommended Dosage in Adults**

33

Concomitant Medications	Dosage of SELZENTRY
Potent CYP3A inhibitors (with or without a potent CYP3A inducer)	150 mg twice daily
including:	
• protease inhibitors (except tipranavir/ritonavir)	
delavirdine	
elvitegravir/ritonavir	
ketoconazole, itraconazole, clarithromycin	
• other potent CYP3A inhibitors (e.g., nefazodone, telithromycin)	
• boceprevir	
Noninteracting concomitant medications, including	300 mg twice daily
tipranavir/ritonavir, nevirapine, raltegravir, all nucleoside reverse	
transcriptase inhibitors (NRTIs), and enfuvirtide <sup>a</sup>	
Potent CYP3A inducers (without a potent CYP3A inhibitor)	600 mg twice daily
including:	
• efavirenz	
• rifampin	
etravirine	
carbamazepine, phenobarbital, and phenytoin	

<sup>a</sup> Noninteracting concomitant medications include all medications that are not potent CYP3A

inhibitors or inducers.

# 36 2.4 Recommended Dosage in Pediatric Patients

- 37 The recommended dosage of SELZENTRY should be based on body weight (kg) and should not
- 38 exceed the recommended adult dose. The recommended dosage also differs based on
- 39 concomitant medications due to drug interactions (Table 2 and Table 3) [see Drug Interactions
- 40 (7.1), Use in Specific Populations (8.4)].
- 41 Before prescribing SELZENTRY tablets, assess children for the ability to swallow tablets. If a
- child is unable to reliably swallow SELZENTRY tablets, the oral solution formulation should be
- prescribed. Administer the oral solution using the included press-in bottle adapter and oral
- 44 dosing syringe.

#### 45 Table 2. Recommended Dosage in Pediatric Patients Aged 2 Years and Older Weighing

#### 46 at Least 10 kg (Tablets)

	Dosage of SELZENTRY Based on Weight					
	10 kg to	10 kg to 20 kg to 30 kg to				
<b>Concomitant Medications</b>	<20 kg	<30 kg	<40 kg	≥40 kg		
Potent CYP3A inhibitors (with	50 mg	75 mg	100 mg	150 mg		
or without a CYP3A inducer)	twice daily	twice daily	twice daily	twice daily		
including:						

enfuvirtide <sup>a</sup> Potent CYP3A inducers (without a potent CYP3A inhibitor) including:  • efavirenz  • rifampin  • etravirine  • carbamazepine,		Not recomi	nended	
medications, including tipranavir/ritonavir, nevirapine, raltegravir, all NRTIs, and	recommended	recommended	twice daily	twice daily
<ul> <li>protease inhibitors (except tipranavir/ritonavir)</li> <li>delavirdine</li> <li>elvitegravir/ritonavir</li> <li>ketoconazole, itraconazole, clarithromycin</li> <li>other potent CYP3A inhibitors (e.g., nefazodone, telithromycin)</li> <li>boceprevir</li> <li>Noninteracting concomitant</li> </ul>	Not	Not	300 mg	300 mg

<sup>&</sup>lt;sup>a</sup> Noninteracting concomitant medications include all medications that are not potent CYP3A

# 49 Table 3. Recommended Dosage in Pediatric Patients Aged 2 Years and Older Weighing

50 at Least 10 kg (Oral Solution)

	Dosage (Volume of Solution) of SELZENTRY						
Concomitant Medications	10 kg to <20 kg						
Potent CYP3A inhibitors (with or without a CYP3A inducer) including:  • protease inhibitors (except tipranavir/ritonavir)  • delavirdine  • elvitegravir/ritonavir  • ketoconazole, itraconazole,	50 mg (2.5 mL) twice daily	<30 kg 80 mg (4 mL) twice daily	<40 kg 100 mg (5 mL) twice daily	≥40 kg 150 mg (7.5 mL) twice daily			

inhibitors or inducers.

<ul> <li>clarithromycin</li> <li>other potent CYP3A inhibitors (e.g., nefazodone, telithromycin)</li> <li>boceprevir</li> </ul>				
Noninteracting concomitant medications, including tipranavir/ritonavir, nevirapine, raltegravir, all NRTIs, and enfuvirtide <sup>a</sup>	Not recommended	Not recommended	300 mg (15 mL) twice daily	300 mg (15 mL) twice daily
Potent CYP3A inducers (without a potent CYP3A inhibitor) including:  • efavirenz  • rifampin  • etravirine  • carbamazepine, phenobarbital, and phenytoin		Not recomm	nended	

<sup>&</sup>lt;sup>a</sup> Noninteracting concomitant medications include all medications that are not potent CYP3A

# **2.5** Recommended Dosage in Patients with Renal Impairment

54 Adults

57

- Table 4 provides dosing recommendations for patients based on renal function and concomitant
- 56 medications.

# Table 4. Recommended Dosage in Adults Based on Renal Function

	Dosage of SELZENTRY Based on Renal Function					
		Mild	Moderate		End-Stage	
	Normal	(CrCl >50	(CrCl≥30	Severe	Renal Disease	
	(CrCl >80	and ≤80	and ≤50	(CrCl <30	on Regular	
<b>Concomitant Medications</b>	mL/min)	mL/min)	mL/min)	mL/min)	Hemodialysis	
Potent CYP3A inhibitors	150 mg	150 mg	150 mg	Contra-	Contra-	
(with or without a CYP3A	twice daily	twice daily	twice daily	indicated	indicated	
inducer) including:						
<ul> <li>protease inhibitors</li> </ul>						
(except						
tipranavir/ritonavir)						
<ul> <li>delavirdine</li> </ul>						
• elvitegravir/ritonavir						

<sup>52</sup> inhibitors or inducers.

<ul> <li>ketoconazole,         itraconazole,         clarithromycin</li> <li>other potent CYP3A         inhibitors (e.g.,</li> </ul>					
nefazodone, telithromycin)					
boceprevir					
Noninteracting concomitant medications including tipranavir/ritonavir,	300 mg twice daily	300 mg twice daily	300 mg twice daily	300 mg twice daily <sup>b</sup>	300 mg twice daily <sup>b</sup>
nevirapine, raltegravir, all					
NRTIs, and enfuvirtide <sup>a</sup>					
Potent CYP3A inducers (without a potent CYP3A inhibitor) including: • efavirenz	600 mg twice daily	600 mg twice daily	600 mg twice daily	Contra- indicated	Contra- indicated
<ul><li>rifampin</li><li>etravirine</li></ul>					
carbamazepine,     phenobarbital, and     phenytoin					

- a Noninteracting concomitant medications include all medications that are not potent CYP3A
   inhibitors or inducers.
- 60 b The dosage of SELZENTRY should be reduced to 150 mg twice daily if there are any
- 61 symptoms of postural hypotension [see Contraindications (4), Warnings and Precautions
- 62 (5.3)].

#### 63 Pediatric Patients

- There are no data to recommend specific doses of SELZENTRY in pediatric patients with mild
- or moderate renal impairment [see Use in Specific Populations (8.6)]. Additionally,
- SELZENTRY is contraindicated for pediatric patients with severe renal impairment or end-stage
- 67 renal disease (ESRD) on regular hemodialysis who are receiving potent CYP3A inhibitors [see
- 68 *Contraindications* (4)].

#### 3 DOSAGE FORMS AND STRENGTHS

70 Tablets:

69

• 25-mg blue, oval, film-coated tablets debossed with "MVC 25" on one side and plain on the other.

- 75-mg blue, oval, film-coated tablets debossed with "MVC 75" on one side and plain on the other.
- 150-mg blue, oval, film-coated tablets debossed with "MVC 150" on one side and plain on the other.
- 300-mg blue, oval, film-coated tablets debossed with "MVC 300" on one side and plain on the other.
- 79 Oral Solution:
- 20 mg per mL clear, colorless, strawberry-flavored oral solution.

#### 81 4 CONTRAINDICATIONS

- 82 SELZENTRY is contraindicated in patients with severe renal impairment or ESRD (CrCl less
- than 30 mL per minute) who are concomitantly taking potent CYP3A inhibitors or inducers [see
- 84 *Warnings and Precautions (5.3)].*

#### 85 5 WARNINGS AND PRECAUTIONS

# 86 **5.1 Hepatotoxicity**

- 87 Hepatotoxicity with allergic features including life-threatening events has been reported in
- 88 clinical trials and postmarketing. Severe rash or evidence of systemic allergic reaction including
- 89 drug-related rash with fever, eosinophilia, elevated IgE, or other systemic symptoms have been
- 90 reported in conjunction with hepatotoxicity [see Warnings and Precautions (5.2)]. These events
- occurred approximately 1 month after starting treatment. Among reported cases of hepatitis,
- some were observed in the absence of allergic features or with no pre-existing hepatic disease.
- Appropriate laboratory testing including ALT, AST, and bilirubin should be conducted prior to
- 94 initiating therapy with SELZENTRY and at other time points during treatment as clinically
- 95 indicated. Hepatic laboratory parameters should be obtained in any patient who develops rash, or
- signs or symptoms of hepatitis, or allergic reaction. Discontinuation of SELZENTRY should be
- onsidered in any patient with signs or symptoms of hepatitis, or with increased liver
- transaminases combined with rash or other systemic symptoms.
- When administering SELZENTRY to patients with pre-existing liver dysfunction or who are co-
- infected with hepatitis B and/or C virus, additional monitoring may be warranted. The safety and
- efficacy of SELZENTRY have not been specifically studied in patients with significant
- underlying liver disorders.

# 103 **5.2** Severe Skin and Hypersensitivity Reactions

- Severe, potentially life-threatening skin and hypersensitivity reactions have been reported in
- patients taking SELZENTRY, in most cases concomitantly with other drugs associated with
- these reactions. These include cases of Stevens-Johnson syndrome (SJS), toxic epidermal

- necrolysis (TEN), and drug rash with eosinophilia and systemic symptoms (DRESS) [see
- 108 Adverse Reactions (6.3)]. The cases were characterized by features including rash, constitutional
- findings, and sometimes organ dysfunction, including hepatic failure. Discontinue SELZENTRY
- and other suspected agents immediately if signs or symptoms of severe skin or hypersensitivity
- reactions develop (including, but not limited to, severe rash or rash accompanied by fever,
- malaise, muscle or joint aches, blisters, oral lesions, conjunctivitis, facial edema, lip swelling,
- eosinophilia). Delay in stopping treatment with SELZENTRY or other suspect drugs after the
- onset of rash may result in a life-threatening reaction. Clinical status, including liver
- aminotransferases, should be monitored and appropriate therapy initiated.

## 116 **5.3** Cardiovascular Events

- Eleven subjects (1.3%) who received SELZENTRY had cardiovascular events, including
- myocardial ischemia and/or infarction, during the Phase 3 trials in treatment-experienced
- subjects (total exposure 609 patient-years [300 on SELZENTRY once daily + 309 on
- 120 SELZENTRY twice daily]), while no subjects who received placebo had such events (total
- exposure 111 patient-years). These subjects generally had cardiac disease or cardiac risk factors
- prior to use of SELZENTRY, and the relative contribution of SELZENTRY to these events is
- 123 not known.
- In the Phase 2b/3 trial in treatment-naive adult subjects, 3 subjects (0.8%) who received
- SELZENTRY had events related to ischemic heart disease and 5 subjects (1.4%) who received
- efavirenz had such events (total exposure 506 and 508 patient-years for SELZENTRY and
- 127 efavirenz, respectively).
- When SELZENTRY was administered to healthy volunteers at doses higher than the
- recommended dose, symptomatic postural hypotension was seen at a greater frequency than in
- placebo. However, when SELZENTRY was given at the recommended dose in HIV-1-infected
- adult subjects in Phase 3 trials, postural hypotension was seen at a rate similar to placebo
- 132 (approximately 0.5%).
- Patients with cardiovascular comorbidities, risk factors for postural hypotension, or receiving
- 134 concomitant medication known to lower blood pressure, could be at increased risk of
- cardiovascular adverse events triggered by postural hypotension. Additional monitoring may be
- warranted.
- 137 <u>Postural Hypotension in Patients with Renal Impairment</u>
- An increased risk of postural hypotension may occur in patients with severe renal insufficiency
- or in those with ESRD due to increased maraviroc exposure in some patients. SELZENTRY
- should be used in patients with severe renal impairment or ESRD only if they are not receiving a
- 141 concomitant potent CYP3A inhibitor or inducer. However, the use of SELZENTRY in these
- patients should only be considered when no alternative treatment options are available. If adult
- patients with severe renal impairment or ESRD experience any symptoms of postural

- 144 hypotension while taking 300 mg twice daily, the dose should be reduced to 150 mg twice daily
- 145 [see Dosage and Administration (2.5)].

# 146 **5.4 Immune Reconstitution Syndrome**

- 147 Immune reconstitution syndrome has been reported in patients treated with combination
- antiretroviral therapy, including SELZENTRY. During the initial phase of combination
- antiretroviral treatment, patients whose immune systems respond may develop an inflammatory
- response to indolent or residual opportunistic infections (such as infection with *Mycobacterium*
- avium infection, cytomegalovirus, Pneumocystis jirovecii pneumonia [PCP], tuberculosis, or
- reactivation of *Herpes simplex* and *Herpes zoster*), which may necessitate further evaluation and
- 153 treatment.
- 154 Autoimmune disorders (such as Graves' disease, polymyositis, and Guillain-Barré syndrome)
- have also been reported to occur in the setting of immune reconstitution; however, the time to
- onset is more variable, and can occur many months after initiation of treatment.

#### 157 **5.5 Potential Risk of Infection**

- SELZENTRY antagonizes the CCR5 co-receptor located on some immune cells, and therefore
- 159 could potentially increase the risk of developing infections. The overall incidence and severity of
- infection, as well as AIDS-defining category C infections, were comparable in the treatment
- groups during the Phase 3 adult treatment-experienced trials of SELZENTRY. While there was a
- higher rate of certain upper respiratory tract infections reported in the treatment arm receiving
- SELZENTRY compared with placebo (23% versus 13%), there was a lower rate of pneumonia
- 164 (2% versus 5%) reported in subjects receiving SELZENTRY. A higher incidence of Herpes virus
- infections (11 per 100 patient-years) was also reported in the treatment arm receiving
- SELZENTRY when adjusted for exposure compared with placebo (8 per 100 patient-years).
- In the Phase 2b/3 trial in treatment-naive adult subjects, the incidence of AIDS-defining
- 168 Category C events when adjusted for exposure was 1.8 for SELZENTRY compared with 2.4 for
- efavirenz per 100 patient-years of exposure.
- 170 Patients should be monitored closely for evidence of infections while receiving SELZENTRY.

# 171 **5.6 Potential Risk of Malignancy**

- While no increase in malignancy has been observed with SELZENTRY, due to this drug's
- mechanism of action, it could affect immune surveillance and lead to an increased risk of
- 174 malignancy.
- 175 The exposure-adjusted rate for malignancies per 100 patient-years of exposure in adult
- treatment-experienced trials was 4.6 for SELZENTRY compared with 9.3 on placebo. In
- treatment-naive adult subjects, the rates were 1.0 and 2.4 per 100 patient-years of exposure for
- 178 SELZENTRY and efavirenz, respectively.
- Long-term follow-up is needed to more fully assess this risk.

#### 180 **6 ADVERSE REACTIONS**

- 181 The following adverse reactions are discussed in other sections of the labeling:
- Hepatotoxicity [see Boxed Warning, Warnings and Precautions (5.1)]
- Severe Skin and Hypersensitivity Reactions [see Warnings and Precautions (5.2)]
- Cardiovascular Events [see Warnings and Precautions (5.3)]

#### 185 **6.1** Clinical Trials Experience

- Because clinical trials are conducted under widely varying conditions, adverse reaction rates
- observed in the clinical trials of a drug cannot be directly compared with rates in the clinical
- trials of another drug and may not reflect the rates observed in practice.
- 189 Adverse Reactions in Adult Subjects
- 190 Treatment-Experienced Subjects: The safety profile of SELZENTRY is primarily based on
- 191 840 HIV-1-infected subjects who received at least 1 dose of SELZENTRY during two Phase 3
- trials. A total of 426 of these subjects received the indicated twice-daily dosing regimen.
- Assessment of treatment-emergent adverse events is based on the pooled data from 2 trials in
- subjects with CCR5-tropic HIV-1 (A4001027 and A4001028). The median duration of therapy
- with SELZENTRY for subjects in these trials was 48 weeks, with the total exposure on
- 196 SELZENTRY twice daily at 309 patient-years versus 111 patient-years on placebo each
- administered with optimized background therapy (OBT). The population was 89% male and
- 198 84% white, with mean age of 46 years (range: 17 to 75 years). Subjects received dose
- 199 equivalents of 300 mg maraviroc once or twice daily.
- The most common adverse events reported with twice-daily therapy with SELZENTRY with
- frequency rates higher than placebo, regardless of causality, were upper respiratory tract
- infections, cough, pyrexia, rash, and dizziness. In these 2 trials, the rate of discontinuation due to
- adverse events was 5% for subjects who received SELZENTRY twice daily + OBT as well as
- 204 those who received placebo + OBT. Most of the adverse events reported were judged to be mild
- 205 to moderate in severity. The data described below occurred with twice-daily dosing of
- 206 SELZENTRY.
- The total numbers of subjects reporting infections were 233 (55%) and 84 (40%) in the group
- 208 receiving SELZENTRY twice daily and the placebo group, respectively. Correcting for the
- 209 longer duration of exposure on SELZENTRY compared with placebo, the exposure-adjusted
- 210 frequency (rate per 100 subject-years) of these events was 133 for both SELZENTRY twice
- daily and placebo.
- 212 Dizziness or postural dizziness occurred in 8% of subjects on either SELZENTRY or placebo,
- with 2 subjects (0.5%) on SELZENTRY permanently discontinuing therapy (1 due to syncope, 1
- due to orthostatic hypotension) versus 1 subject on placebo (0.5%) permanently discontinuing
- 215 therapy due to dizziness.

Treatment-emergent adverse events, regardless of causality, from Trials A4001027 and A4001028 are summarized in Table 5. Selected events occurring at greater than or equal to 2% of subjects and at a numerically higher rate in subjects treated with SELZENTRY are included; events that occurred at the same or higher rate on placebo are not displayed.

Table 5. Selected Treatment-Emergent Adverse Events (All Causality) ≥2% on
 SELZENTRY (and at a Higher Rate Compared with Placebo) in Trials A4001027 and
 A4001028 (Pooled Analysis, 48 Weeks)

	SELZENTRY			
	Twice Daily <sup>a</sup>			Placebo
		Exposure-		Exposure-
		Adjusted Rate		Adjusted Rate
Body System/	(n = 426)	(per 100 pt-yrs)	(n = 209)	(per 100 pt-yrs)
Adverse Event	%	$PYE = 309^{b}$	%	$\mathbf{PYE} = 111^{\mathbf{b}}$
Eye Disorders				
Conjunctivitis	2	3	1	3
Ocular infections, inflammations,	2	3	1	2
and associated manifestations				
Gastrointestinal Disorders				
Constipation	6	9	3	6
General Disorders and				
Administration Site Conditions				
Pyrexia	13	20	9	17
Pain and discomfort	4	5	3	5
Infections and Infestations				
Upper respiratory tract infection	23	37	13	27
Herpes infection	8	11	4	8
Sinusitis	7	10	3	6
Bronchitis	7	9	5	9
Folliculitis	4	5	2	4
Anogenital warts	2	3	1	3
Influenza	2	3	0.5	1
Otitis media	2	3	0.5	1
Metabolism and Nutrition				
Disorders				
Appetite disorders	8	11	7	13
Musculoskeletal and Connective				
Tissue Disorders				
Joint-related signs and symptoms	7	10	3	5
Muscle pains	3	4	0.5	1

Neoplasms Benign, Malignant,				
and Unspecified				
Skin neoplasms benign	3	4	1	3
Nervous System Disorders				
Dizziness/postural dizziness	9	13	8	17
Paresthesias and dysesthesias	5	7	3	6
Sensory abnormalities	4	6	1	3
Disturbances in consciousness	4	5	3	6
Peripheral neuropathies	4	5	3	6
Psychiatric Disorders				
Disturbances in initiating and	8	11	5	10
maintaining sleep				
Depressive disorders	4	6	3	5
Anxiety symptoms	4	5	3	7
Renal and Urinary Disorders				
Bladder and urethral symptoms	5	7	1	3
Urinary tract signs and symptoms	3	4	1	3
Respiratory, Thoracic, and				
Mediastinal Disorders				
Coughing and associated	14	21	5	10
symptoms				
Upper respiratory tract signs and	6	9	3	6
symptoms				
Nasal congestion and	4	6	3	5
inflammations				
Breathing abnormalities	4	5	2	5
Paranasal sinus disorders	3	4	0.5	1
Skin and Subcutaneous Tissue				
Disorders				
Rash	11	16	5	11
Apocrine and eccrine gland	5	7	4	7.5
disorders				
Pruritus	4	5	2	4
Lipodystrophies	3	5	0.5	1
Erythema	2	3	1	2
Vascular Disorders				
Vascular hypertensive disorders	3	4	2	4

 <sup>&</sup>lt;sup>a</sup> 300-mg dose equivalent.
 <sup>b</sup> PYE = Patient-years of exposure. 

225 Laboratory Abnormalities: Table 6 shows the treatment-emergent Grade 3-4 laboratory

abnormalities that occurred in greater than 2% of subjects receiving SELZENTRY.

**Table 6. Maximum Shift in Laboratory Test Values (without Regard to Baseline) ≥2%** 

of Grade 3-4 Abnormalities (ACTG Criteria) in Trials A4001027 and A4001028 (Pooled

229 Analysis, 48 Weeks)

227

228

Laboratory Parameter Preferred Term	Limit	SELZENTRY Twice Daily + OBT (n = 421) <sup>a</sup>	Placebo + OBT (n = 207) <sup>a</sup>
Aspartate aminotransferase	>5.0 x ULN	4.8	2.9
Alanine aminotransferase	>5.0 x ULN	2.6	3.4
Total bilirubin	>2.5 x ULN	5.5	5.3
Amylase	>2.0 x ULN	5.7	5.8
Lipase	>2.0 x ULN	4.9	6.3
Absolute neutrophil count	<750/mm <sup>3</sup>	4.3	2.4

230 ULN = Upper limit of normal.

<sup>a</sup> Percentages based on total subjects evaluated for each laboratory parameter.

232 Treatment-Naive Subjects: Treatment-Emergent Adverse Events: Treatment-emergent adverse

events, regardless of causality, from Trial A4001026, a double-blind, comparative, controlled

234 trial in which 721 treatment-naive subjects received SELZENTRY 300 mg twice daily (n = 360)

or efavirenz 600 mg once daily (n = 361) in combination with lamivudine/zidovudine

236 (COMBIVIR) for 96 weeks, are summarized in Table 7. Selected events occurring in greater

than or equal to 2% of subjects and at a numerically higher rate in subjects treated with

SELZENTRY are included; events that occurred at the same or higher rate on efavirenz are not

239 displayed.

Table 7. Selected Treatment-Emergent Adverse Events (All Causality) ≥2% on

241 SELZENTRY (and at a Higher Rate Compared with Efavirenz) in Trial A4001026 (96

242 Weeks)

240

	SELZENTRY	Efavirenz
	300 mg Twice Daily +	600 mg Once Daily +
	Lamivudine/Zidovudine	Lamivudine/Zidovudine
Body System/	(n = 360)	(n = 361)
Adverse Event	%	%
Blood and Lymphatic System		
Disorders		
Anemias NEC	8	5
Neutropenias	4	3
Ear and Labyrinth Disorders		

Ear disorders NEC	3	2
<b>Gastrointestinal Disorders</b>		
Flatulence, bloating, and distention	10	7
Gastrointestinal atonic and	9	5
hypomotility disorders NEC		
Gastrointestinal signs and	3	2
symptoms NEC		
General Disorders and		
<b>Administration Site Conditions</b>		
Body temperature perception	3	1
Infections and Infestations		
Upper respiratory tract infection	32	30
Bronchitis	13	9
Herpes infection	7	6
Bacterial infections NEC	6	3
Herpes zoster/varicella	5	4
Tinea infections	4	3
Lower respiratory tract and lung	3	2
infections		
Neisseria infections	3	0
Viral infections NEC	3	2
Musculoskeletal and Connective		
Tissue Disorders		
Joint-related signs and symptoms	6	5
Nervous System Disorders		
Paresthesias and dysesthesias	4	3
Memory loss (excluding dementia)	3	1
Renal and Urinary Disorders		
Bladder and urethral symptoms	4	3
Reproductive System and Breast		
Disorders		
Erection and ejaculation conditions	3	2
and disorders		
Respiratory, Thoracic, and		
Mediastinal Disorders		
Upper respiratory tract signs and	9	5
symptoms		
Skin and Subcutaneous Disorders		
Nail and nail bed conditions	6	2

(excluding infections and		
infestations)		
Lipodystrophies	4	3
Acnes	3	2
Alopecias	2	1

243 Laboratory Abnormalities:

Table 8. Maximum Shift in Laboratory Test Values (without Regard to Baseline) ≥2% of Grade 3-4 Abnormalities (ACTG Criteria) in Trial A4001026 (96 Weeks)

Laboratory		SELZENTRY 300 mg Twice Daily + Lamivudine/Zidovudine	Efavirenz 600 mg Once Daily+ Lamiyudine/Zidoyudine
Parameter		$(n = 353)^a$	$(n = 350)^a$
Preferred Term	Limit	%	0/0
Aspartate	>5.0 x ULN	4.0	4.0
aminotransferase			
Alanine	>5.0 x ULN	3.9	4.0
aminotransferase			
Creatine kinase	>10.0 x ULN	3.9	4.8
Amylase	>2.0 x ULN	4.3	6.0
Absolute neutrophil	<750/mm <sup>3</sup>	5.7	4.9
count			
Hemoglobin	<7.0 g/dL	2.9	2.3

<sup>246</sup> ULN = Upper limit of normal.

in a given treatment group had greater than 1 occurrence of the same abnormality, only the

250 most severe is counted.

248

249

253

254

256

257

258

259

260

251 Less Common Adverse Events in Clinical Trials: The following adverse events occurred in less

252 than 2% of subjects treated with SELZENTRY or at a rate similar to the comparator. These

events have been included because of their seriousness and either increased frequency on

SELZENTRY or are potential risks due to the mechanism of action. Events attributed to the

subjects' underlying HIV-1 infection are not listed.

Blood and Lymphatic System: Marrow depression and hypoplastic anemia.

Cardiac Disorders: Unstable angina, acute cardiac failure, coronary artery disease,

coronary artery occlusion, myocardial infarction, myocardial ischemia.

*Hepatobiliary Disorders:* Hepatic cirrhosis, hepatic failure, cholestatic jaundice, portal vein thrombosis, jaundice.

-

<sup>&</sup>lt;sup>a</sup> n = Total number of subjects evaluable for laboratory abnormalities.

Percentages based on total subjects evaluated for each laboratory parameter. If the same subject

261 262	<i>Infections and Infestations:</i> Endocarditis, infective myositis, viral meningitis, pneumonia, treponema infections, septic shock, <i>Clostridium</i> difficile colitis, meningitis.
<ul><li>263</li><li>264</li></ul>	Musculoskeletal and Connective Tissue Disorders: Myositis, osteonecrosis, rhabdomyolysis, blood CK increased.
265 266 267 268 269 270	Neoplasms Benign, Malignant, and Unspecified (Including Cysts and Polyps): Abdominal neoplasm, anal cancer, basal cell carcinoma, Bowen's disease, cholangiocarcinoma, diffuse large B-cell lymphoma, lymphoma, metastases to liver, esophageal carcinoma, nasopharyngeal carcinoma, squamous cell carcinoma, squamous cell carcinoma of skin, tongue neoplasm (malignant stage unspecified), anaplastic large cell lymphomas T- and null-cell types, bile duct neoplasms malignant, endocrine neoplasms malignant and unspecified.
271 272	Nervous System Disorders: Cerebrovascular accident, convulsions and epilepsy, tremor (excluding congenital), facial palsy, hemianopia, loss of consciousness, visual field defect.
273	Clinical Trials Experience in Pediatric Subjects
274 275 276 277 278 279 280	Trial A4001031 is an open-label trial in which 103 treatment-experienced, CCR5-tropic, HIV-1–infected pediatric subjects aged 2 to less than 18 years weighing at least 10 kg received SELZENTRY twice daily in combination with OBT. The dose of SELZENTRY was based on body surface area (BSA) and on whether the subject was receiving potent CYP3A inhibitors and/or inducers. The median duration of therapy with SELZENTRY was 131 weeks with 72% of subjects receiving study treatment for greater than 48 weeks and 62% of subjects receiving study treatment for 96 weeks.
281 282 283 284 285 286	In these 103 children and adolescents, the safety profile through 96 weeks was similar to that for adults. Most of the adverse reactions reported were mild to moderate; severe (Grade 3 and 4) adverse reactions occurred in 2% of subjects. The most common adverse reactions (all grades) reported with twice-daily therapy with SELZENTRY were vomiting (12%), abdominal pain (4%), diarrhea (4%), nausea (4%), and dizziness (3%). Three subjects (3%) discontinued due to adverse events.
287 288 289 290	Maraviroc-related gastrointestinal adverse events through 48 weeks (nausea, vomiting, diarrhea, constipation, and abdominal pain/cramps) were observed more commonly in subjects who received the SELZENTRY oral solution (21%) compared with those who received SELZENTRY tablets (16%). Subjects were permitted to change formulations after Week 48.
291	6.2 Postmarketing Experience
<ul><li>292</li><li>293</li><li>294</li></ul>	The following adverse events have been identified during post-approval use of SELZENTRY. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug

# Skin and Subcutaneous Tissue Disorders

295

296

exposure.

- 297 Stevens-Johnson syndrome (SJS), drug rash with eosinophilia and systemic symptoms (DRESS),
- 298 toxic epidermal necrolysis (TEN).

#### 299 **7 DRUG INTERACTIONS**

# 300 7.1 Effect of Concomitant Drugs on the Pharmacokinetics of Maraviroc

- 301 Maraviroc is metabolized by CYP3A, and is also a substrate for P-glycoprotein (P-gp), organic
- anion-transporting polypeptide (OATP)1B1, and multidrug resistance-associated protein
- 303 (MRP)2. The pharmacokinetics of maraviroc are likely to be modulated by inhibitors and
- inducers of CYP3A and P-gp, and may be modulated by inhibitors of OATP1B1 and MRP2.
- Therefore, a dosage adjustment may be required when maraviroc is coadministered with those
- 306 drugs [see Dosage and Administration (2.3, 2.4)].
- 307 Concomitant use of maraviroc and St. John's wort (*Hypericum perforatum*) or products
- 308 containing St. John's wort is not recommended. Coadministration of maraviroc with St. John's
- wort is expected to substantially decrease maraviroc concentrations and may result in suboptimal
- 310 levels of maraviroc and lead to loss of virologic response and possible resistance to maraviroc.
- 311 Additional drug interaction information is available [see *Clinical Pharmacology (12.3)*].

#### 312 8 USE IN SPECIFIC POPULATIONS

# 313 **8.1 Pregnancy**

- 314 Pregnancy Exposure Registry
- There is a pregnancy exposure registry that monitors pregnancy outcomes in women exposed to
- 316 SELZENTRY during pregnancy. Physicians are encouraged to register patients by calling the
- 317 Antiretroviral Pregnancy Registry (APR) at 1-800-258-4263.
- 318 Risk Summary
- Limited data on the use of SELZENTRY during pregnancy from the APR and case reports are
- 320 not sufficient to inform a drug-associated risk of birth defects and miscarriage. In animal
- 321 reproduction studies, no evidence of adverse developmental outcomes was observed with
- maraviroc. During organogenesis in the rat and rabbit, systemic exposures (AUC) to maraviroc
- were approximately 20 times (rats) and 5 times (rabbits) the exposure in humans at the
- recommended 300-mg twice-daily dose. In the rat pre- and post-natal development study,
- maternal systemic exposure (AUC) to maraviroc was approximately 14 times the exposure in
- humans at the recommended 300-mg twice-daily dose (see Data).
- 327 The estimated background risk of major birth defects and miscarriage for the indicated
- 328 population is unknown. All pregnancies have a background risk of birth defect, loss, or other
- adverse outcomes. In the U.S. general population, the estimated background risk of major birth
- defects and miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%,
- 331 respectively.

332	Data

- 333 Animal Data: Maraviroc was administered orally to pregnant rats (up to 1,000 mg per kg per
- day) and rabbits (up to 75 mg per kg per day) on gestation Days 6 to 17 and 7 to 19, respectively.
- No adverse effects on embryo-fetal development were observed at these dose levels, resulting in
- exposures (AUC) approximately 20 times (rats) and 5 times (rabbits) higher than human
- exposures at the recommended daily dose. In the rat pre- and post-natal development study,
- maraviroc was administered orally at up to 1,000 mg per kg per day on gestation Day 6 to
- lactation/post-partum Day 20, with development of the offspring (including fertility and
- 340 reproductive performance) unaffected by maternal administration of maraviroc at an exposure
- 341 (AUC) approximately 14 times higher than human exposure at the recommended daily dose.

#### 342 **8.2** Lactation

# 343 Risk Summary

- 344 The Centers for Disease Control and Prevention recommend that HIV-1-infected mothers in the
- 345 United States not breastfeed their infants to avoid risking postnatal transmission of HIV-1
- 346 infection.
- 347 There are no data on the presence of maraviroc in human milk, the effects on the breastfed
- infant, or the effects on milk production. When administered to lactating rats, maraviroc was
- present in milk (see Data). Because of the potential for (1) HIV transmission (in HIV-negative
- infants), (2) developing viral resistance (in HIV-positive infants), and (3) serious adverse
- reactions in a breastfed infant similar to those seen in adults, instruct mothers not to breastfeed if
- 352 they are receiving SELZENTRY.
- 353 Data
- Maraviroc (and related metabolites) was excreted into the milk of lactating rats following a
- single oral dose of maraviroc (100 mg per kg) on lactation Day 12, with a maximal milk
- concentration achieved one hour post-administration at a milk concentration approximately 2.5
- 357 times that of maternal plasma concentrations.

#### 358 **8.4 Pediatric Use**

- 359 The safety, pharmacokinetic (PK) profile, and antiviral activity of SELZENTRY were evaluated
- in treatment-experienced, CCR5-tropic, HIV-1-infected pediatric subjects aged 2 to less than 18
- years weighing at least 10 kg in an open-label, multicenter clinical trial, A4001031 [see Adverse]
- 362 Reactions (6.1), Clinical Studies (14.2)]. Pharmacokinetics were evaluated in a total of 98
- 363 pediatric subjects: 85 subjects received SELZENTRY and concomitant medications that included
- potent CYP3A inhibitors with or without potent CYP3A inducers, 10 subjects received
- 365 SELZENTRY and noninteracting medications (not containing potent CYP3A inhibitors or potent
- 366 CYP3A inducers), and three subjects received SELZENTRY and medications that included
- potent CYP3A inducers without potent CYP3A inhibitors [see Clinical Pharmacology (12.3)].

- 368 See *Dosage and Administration* (2.4, 2.5) for dosing recommendations for pediatric patients
- aged 2 years and older and weighing at least 10 kg. The pharmacokinetics, safety, and efficacy of
- maraviroc in patients younger than 2 years have not been established. Therefore, SELZENTRY
- is not recommended in this patient population. Additionally, there are insufficient data to make
- dosing recommendations for use of SELZENTRY in pediatric patients concomitantly receiving
- 373 noninteracting medications and weighing less than 30 kg or in pediatric patients concomitantly
- 374 receiving potent CYP3A inducers without a potent CYP3A inhibitor [see Dosage and
- 375 *Administration* (2.4, 2.5)].

#### 8.5 Geriatric Use

376

- 377 There were insufficient numbers of subjects aged 65 and over in the clinical trials to determine
- whether they respond differently from younger subjects. In general, caution should be exercised
- 379 when administering SELZENTRY in elderly patients, also reflecting the greater frequency of
- decreased hepatic and renal function, of concomitant disease and other drug therapy.

# 381 **8.6 Renal Impairment**

- 382 Recommended doses of SELZENTRY for adult patients with impaired renal function (CrCl less
- than or equal to 80 mL per minute) are based on the results of a pharmacokinetic trial conducted
- in healthy adult subjects with various degrees of renal impairment. Maraviroc has not been
- studied in pediatric patients with renal impairment. There are no data to recommend specific
- doses of SELZENTRY in pediatric patients with mild to moderate renal impairment [see Use in
- 387 Specific Populations (8.4)]. SELZENTRY is contraindicated in pediatric patients with severe
- renal impairment or ESRD on regular hemodialysis who are receiving potent CYP3A inhibitors
- 389 [see Contraindications (4)].
- 390 The pharmacokinetics of maraviroc in adult subjects with mild and moderate renal impairment
- was similar to that in subjects with normal renal function [see Clinical Pharmacology (12.3)]. A
- 392 limited number of adult subjects with mild and moderate renal impairment in the Phase 3 clinical
- trials (n = 131 and n = 12, respectively) received the same dose of SELZENTRY as that
- 394 administered to subjects with normal renal function. In these subjects, there was no apparent
- difference in the adverse event profile for maraviroc compared with subjects with normal renal
- 396 function.
- 397 If adult patients with severe renal impairment or ESRD not receiving a concomitant potent
- 398 CYP3A inhibitor or inducer experience any symptoms of postural hypotension while taking
- 399 SELZENTRY 300 mg twice daily, the dose should be reduced to 150 mg twice daily. No trials
- 400 have been performed in subjects with severe renal impairment or ESRD co-treated with potent
- 401 CYP3A inhibitors or inducers. Hence, no dose of SELZENTRY can be recommended, and
- 402 SELZENTRY is contraindicated for these patients [see Dosage and Administration (2.3),
- 403 Contraindications (4), Warnings and Precautions (5.3), Clinical Pharmacology (12.3)].

#### 8.7 Hepatic Impairment

404

- 405 Maraviroc is principally metabolized by the liver; therefore, when administering this drug to
- 406 patients with hepatic impairment, maraviroc concentrations may be increased. Maraviroc
- 407 concentrations are higher when SELZENTRY 150 mg is administered with a potent CYP3A
- inhibitor compared with following administration of 300 mg without a CYP3A inhibitor, so
- patients with moderate hepatic impairment who receive SELZENTRY 150 mg with a potent
- 410 CYP3A inhibitor should be monitored closely for maraviroc-associated adverse events.
- 411 Maraviroc has not been studied in subjects with severe hepatic impairment or in pediatric
- 412 patients with any degree of hepatic impairment [see Warnings and Precautions (5.1), Clinical
- 413 *Pharmacology* (12.3)].

#### 414 **10 OVERDOSAGE**

- The highest single dose administered in clinical trials was 1,200 mg. The dose-limiting adverse
- event was postural hypotension, which was observed at 600 mg. While the recommended dose
- 417 for SELZENTRY in patients receiving a CYP3A inducer without a CYP3A inhibitor is 600 mg
- 418 twice daily, this dose is appropriate due to enhanced metabolism.
- 419 Prolongation of the QT interval was seen in dogs and monkeys at plasma concentrations 6 and
- 420 12 times, respectively, those expected in humans at the intended exposure of 300-mg equivalents
- 421 twice daily. However, no significant QT prolongation was seen in the trials in treatment-
- 422 experienced subjects with HIV using the recommended doses of maraviroc, or in a specific
- pharmacokinetic trial to evaluate the potential of maraviroc to prolong the QT interval [see
- 424 Clinical Pharmacology (12.2)].
- There is no specific antidote for overdose with maraviroc. Treatment of overdose should consist
- of general supportive measures including keeping the patient in a supine position, careful
- assessment of patient vital signs, blood pressure, and ECG.
- 428 Administration of activated charcoal may also be used to aid in removal of unabsorbed drug.
- Hemodialysis had a minimal effect on maraviroc clearance and exposure in a trial in subjects
- with ESRD [see Clinical Pharmacology (12.3)].

#### 11 DESCRIPTION

431

- 432 SELZENTRY (maraviroc) is a selective, slowly reversible, small molecule antagonist of the
- interaction between human CCR5 and HIV-1 gp120. Blocking this interaction prevents
- 434 CCR5-tropic HIV-1 entry into cells.
- 435 SELZENTRY film-coated tablets for oral administration contain 25, 75, 150, or 300 mg of
- maraviroc and the following inactive ingredients: dibasic calcium phosphate (anhydrous),
- 437 magnesium stearate, microcrystalline cellulose, and sodium starch glycolate. The film coat
- 438 (Opadry II Blue [85G20583]) contains FD&C blue #2 aluminum lake, soya lecithin,
- polyethylene glycol (macrogol 3350), polyvinyl alcohol, talc, and titanium dioxide.

- 440 SELZENTRY oral solution contains 20 mg per mL of maraviroc and the following inactive
- ingredients: citric acid (anhydrous), purified water, sodium benzoate, sodium citrate dihydrate,
- strawberry flavoring (501440T), and sucralose.
- 443 Maraviroc is chemically described as 4,4-difluoro-*N*-{(1*S*)-3-[*exo*-3-(3-isopropyl-5-methyl-4*H*-
- 444 1,2,4-triazol-4-yl)-8-azabicyclo[3.2.1]oct-8-yl]-1-phenylpropyl}cyclohexanecarboxamide.
- The molecular formula is  $C_{29}H_{41}F_2N_5O$  and the structural formula is:

446

- Maraviroc is a white to pale-colored powder with a molecular weight of 513.67. It is highly
- soluble across the physiological pH range (pH 1.0 to 7.5).

# 449 12 CLINICAL PHARMACOLOGY

- 450 **12.1 Mechanism of Action**
- 451 Maraviroc is an HIV-1 antiviral drug [see Microbiology (12.4)].
- 452 **12.2 Pharmacodynamics**
- 453 Exposure-Response Relationship in Treatment-Experienced Adult Subjects
- The relationship between maraviroc, modeled plasma trough concentration (C<sub>min</sub>) (1 to 9 samples
- per subject taken on up to 7 visits), and virologic response was evaluated in
- 456 973 treatment-experienced HIV-1-infected subjects with varied optimized background
- antiretroviral regimens in Trials A4001027 and A4001028. The C<sub>min</sub>, baseline viral load, baseline
- 458 CD4+ cell count, and overall sensitivity score (OSS) were found to be important predictors of
- virologic success (defined as viral load less than 400 copies per mL at 24 weeks). Table 9
- illustrates the proportions of subjects with virologic success (%) within each C<sub>min</sub> quartile for
- 461 150-mg twice-daily and 300-mg twice-daily groups.

# Table 9. Treatment-Experienced Subjects with Virologic Success by C<sub>min</sub> Quartile (Q1-O4)

<b>~</b> -)	150 mg Twice Daily				300 mg Twice Daily			
		(with CYP3A Inhibitors)			(without CYP3A Inhibitors)			
		Median % Subjects with			Median	% Subjects with		
	n	Cmin	Virologic Success	n	Cmin	Virologic Success		
Placebo	160	-	30.6	35	-	28.6		
Q1	78	33	52.6	22	13	50.0		
Q2	77	87	63.6	22	29	68.2		
Q3	78	166	78.2	22	46	63.6		
Q4	78	279	74.4	22	97	68.2		

### 464 Exposure-Response Relationship in Treatment-Naive Adult Subjects

- The relationship between maraviroc, modeled plasma trough concentration ( $C_{min}$ ) (1 to
- 466 12 samples per subject taken on up to 8 visits), and virologic response was evaluated in
- 467 294 treatment-naive HIV-1-infected subjects receiving maraviroc 300 mg twice daily in
- 468 combination with lamivudine/zidovudine in Trial A4001026. Table 10 illustrates the proportion
- 469 (%) of subjects with virologic success less than 50 copies per mL at 48 weeks within each C<sub>min</sub>
- 470 quartile for the 300-mg twice-daily dose.

462

463

471 Table 10. Treatment-Naive Subjects with Virologic Success by C<sub>min</sub> Quartile (Q1-Q4)

		300 mg Twice Daily						
	n	Median C <sub>min</sub> % Subjects with Virologic So						
Q1	75	23	57.3					
Q2	72	39	72.2					
Q3	73	56	74.0					
Q4	74	81	83.8					

- Eighteen of 75 (24%) subjects in Q1 had no measurable maraviroc concentration on at least one
- occasion versus 1 of 73 and 1 of 74 in Q3 and Q4, respectively.

#### 474 Effects on Electrocardiogram

- 475 A placebo-controlled, randomized, crossover trial to evaluate the effect on the QT interval of
- healthy male and female volunteers was conducted with 3 single oral doses of maraviroc and
- 477 moxifloxacin. The placebo-adjusted mean maximum (upper 1-sided 95% CI) increases in QTc
- 478 from baseline after 100, 300, and 900 mg of maraviroc were -2 (0), -1 (1), and 1 (3) msec,
- 479 respectively, and 13 (15) msec for moxifloxacin 400 mg. No subject in any group had an
- increase in QTc of greater than or equal to 60 msec from baseline. No subject experienced an
- interval exceeding the potentially clinically relevant threshold of 500 msec.

#### 12.3 Pharmacokinetics

### Table 11. Mean Maraviroc Pharmacokinetic Parameters in Adults

Patient Population	Maraviroc Dose	n	AUC <sub>12</sub> (ng.h/mL)	C <sub>max</sub> (ng/mL)	C <sub>min</sub> (ng/mL)
Healthy volunteers	300 mg twice daily	64	2,908	888	43.1
(Phase 1)	300 mg twice daily		2,500	000	13.1
Asymptomatic HIV	300 mg twice daily	8	2,550	618	33.6
subjects (Phase 2a)					
Treatment-experienced	300 mg twice daily	94	1,513	266	37.2
HIV subjects (Phase 3) <sup>a</sup>	150 mg twice daily	375	2,463	332	101
	(+ CYP3A inhibitor)				
Treatment-naive HIV	300 mg twice daily	344	1,865	287	60
subjects (Phase 2b/3) <sup>a</sup>					

- <sup>a</sup> The estimated exposure is lower compared with other trials possibly due to sparse sampling,
- food effect, compliance, and concomitant medications.

# 486 Absorption

482

483

- Peak maraviroc plasma concentrations are attained 0.5 to 4 hours following single oral doses of 1
- 488 to 1,200 mg administered to uninfected volunteers. The pharmacokinetics of oral maraviroc are
- and not dose proportional over the dose range.
- The absolute bioavailability of a 100-mg dose is 23% and is predicted to be 33% at 300 mg.
- 491 Maraviroc is a substrate for the efflux transporter P-gp.
- 492 Effect of Food on Oral Absorption: Coadministration of a 300-mg tablet with a high-fat breakfast
- 493 reduced maraviroc C<sub>max</sub> and AUC by 33% and coadministration of 75 mg of oral solution with a
- high-fat breakfast reduced maraviroc AUC by 73% in healthy adult volunteers. Studies with the
- 495 tablet formulation demonstrated a reduced food effect at higher doses.
- There were no food restrictions in the adult trials (using the tablet formulation) or in the pediatric
- 497 trial (using both tablet and oral solution formulations) that demonstrated the efficacy/antiviral
- 498 activity and safety of maraviroc [see Clinical Studies (14.1, 14.2)].

# 499 <u>Distribution</u>

- Maraviroc is bound (approximately 76%) to human plasma proteins, and shows moderate
- affinity for albumin and alpha-1 acid glycoprotein. The volume of distribution of maraviroc is
- approximately 194 L.

#### 503 Elimination

- 504 *Metabolism:* Trials in humans and in vitro studies using human liver microsomes and expressed
- enzymes have demonstrated that maraviroc is principally metabolized by the cytochrome P450
- system to metabolites that are essentially inactive against HIV-1. In vitro studies indicate that
- 507 CYP3A is the major enzyme responsible for maraviroc metabolism. In vitro studies also indicate

- that polymorphic enzymes CYP2C9, CYP2D6, and CYP2C19 do not contribute significantly to
- the metabolism of maraviroc.
- Maraviroc is the major circulating component (~42% drug-related radioactivity) following a
- single oral dose of 300 mg [<sup>14</sup>C]-maraviroc. The most significant circulating metabolite in
- 512 humans is a secondary amine (~22% radioactivity) formed by N-dealkylation. This polar
- 513 metabolite has no significant pharmacological activity. Other metabolites are products of
- mono-oxidation and are only minor components of plasma drug-related radioactivity.
- 515 Excretion: The terminal half-life of maraviroc following oral dosing to steady state in healthy
- subjects was 14 to 18 hours. A mass balance/excretion trial was conducted using a single 300-mg
- dose of <sup>14</sup>C-labeled maraviroc. Approximately 20% of the radiolabel was recovered in the urine
- and 76% was recovered in the feces over 168 hours. Maraviroc was the major component present
- in urine (mean of 8% dose) and feces (mean of 25% dose). The remainder was excreted as
- 520 metabolites.

# 521 Specific Populations

- 522 Patients with Hepatic Impairment: Maraviroc is primarily metabolized and eliminated by the
- liver. A trial compared the pharmacokinetics of a single 300-mg dose of SELZENTRY in
- subjects with mild (Child-Pugh Class A, n = 8) and moderate (Child-Pugh Class B, n = 8)
- hepatic impairment with pharmacokinetics in healthy subjects (n = 8). The mean  $C_{max}$  and AUC
- were 11% and 25% higher, respectively, for subjects with mild hepatic impairment, and 32% and
- 527 46% higher, respectively, for subjects with moderate hepatic impairment compared with subjects
- with normal hepatic function. These changes do not warrant a dose adjustment. Maraviroc
- 529 concentrations are higher when SELZENTRY 150 mg is administered with a potent CYP3A
- inhibitor compared with following administration of 300 mg without a CYP3A inhibitor, so
- patients with moderate hepatic impairment who receive SELZENTRY 150 mg with a potent
- 532 CYP3A inhibitor should be monitored closely for maraviroc-associated adverse events. The
- 533 pharmacokinetics of maraviroc have not been studied in subjects with severe hepatic impairment
- 534 [see Warnings and Precautions (5.1)].
- Patients with Renal Impairment: A trial compared the pharmacokinetics of a single 300-mg dose
- of SELZENTRY in adult subjects with severe renal impairment (CrCl less than 30 mL per
- minute, n = 6) and ESRD (n = 6) with healthy volunteers (n = 6). Geometric mean ratios for
- 538 maraviroc C<sub>max</sub> and AUC<sub>inf</sub> were 2.4-fold and 3.2-fold higher, respectively, for subjects with
- severe renal impairment, and 1.7-fold and 2.0-fold higher, respectively, for subjects with ESRD
- as compared with subjects with normal renal function in this trial. Hemodialysis had a minimal
- effect on maraviroc clearance and exposure in subjects with ESRD. Exposures observed in
- subjects with severe renal impairment and ESRD were within the range observed in previous
- 300-mg single-dose trials of SELZENTRY in healthy volunteers with normal renal function.
- However, maraviroc exposures in the subjects with normal renal function in this trial were 50%
- lower than those observed in previous trials. Based on the results of this trial, no dose adjustment

- is recommended for patients with renal impairment receiving SELZENTRY without a potent
- 547 CYP3A inhibitor or inducer. However, if patients with severe renal impairment or ESRD
- 548 experience any symptoms of postural hypotension while taking SELZENTRY 300 mg twice
- daily, their dose should be reduced to 150 mg twice daily [see Dosage and Administration (2.3),
- 550 *Warnings and Precautions (5.3)*].
- In addition, the trial compared the pharmacokinetics of multiple-dose SELZENTRY in
- combination with saquinavir/ritonavir 1,000/100 mg twice daily (a potent CYP3A inhibitor
- combination) for 7 days in subjects with mild renal impairment (CrCl greater than 50 and less
- than or equal to 80 mL per minute, n = 6) and moderate renal impairment (CrCl greater than or
- equal to 30 and less than or equal to 50 mL per minute, n = 6) with healthy volunteers with
- normal renal function (n = 6). Subjects received 150 mg of SELZENTRY at different dose
- frequencies (healthy volunteers every 12 hours; mild renal impairment every 24 hours;
- 558 moderate renal impairment every 48 hours). Compared with healthy volunteers (dosed every
- 12 hours), geometric mean ratios for maraviroc AUC<sub>tau</sub>, C<sub>max</sub>, and C<sub>min</sub> were 50% higher, 20%
- higher, and 43% lower, respectively, for subjects with mild renal impairment (dosed every
- 561 24 hours). Geometric mean ratios for maraviroc AUC<sub>tau</sub>, C<sub>max</sub>, and C<sub>min</sub> were 16% higher, 29%
- lower, and 85% lower, respectively, for subjects with moderate renal impairment (dosed every
- 48 hours) compared with healthy volunteers (dosed every 12 hours). Based on the data from this
- trial, no adjustment in dose is recommended for patients with mild or moderate renal impairment
- 565 [see Dosage and Administration (2.3)].
- 566 Pediatric Patients: The pharmacokinetics of maraviroc were evaluated in CCR5-tropic, HIV-1–
- infected, treatment-experienced pediatric subjects aged 2 to less than 18 years. In the dose-
- 568 finding stage of Trial A4001031, doses were administered with food on intensive PK evaluation
- days and optimized to achieve an average concentration over the dosing interval (Cavg) of
- greater than 100 ng per mL. Throughout the trial, on non-intensive PK evaluation days maraviroc
- was taken with or without food. The initial dose of maraviroc was based on BSA and
- 572 concomitant medication category (i.e., presence of CYP3A inhibitors and/or inducers). The
- 573 conversion of dosing to a weight (kg)-band basis in children provides comparable exposures with
- those observed in the trial at the corresponding BSA.
- Maraviroc pharmacokinetic parameters in pediatric subjects receiving potent CYP3A inhibitors
- with or without a potent CYP3A inducer (Table 12) and in subjects weighing greater than or
- equal to 30 kg and receiving noninteracting concomitant medications (Table 13) were similar to
- 578 those observed in adults. Insufficient pharmacokinetic data are available to make a comparison
- between adults and pediatric subjects weighing less than 30 kg and receiving noninteracting
- 580 concomitant medications or between adult and pediatric subjects receiving concomitant
- medications consisting of a potent CYP3A inducer without CYP3A inhibitor.

585

586

587

588

589

590

600

		Maraviroc Pharmacokinetic Parameter <sup>a</sup> Geometric Mean				
Weight	Dose of SELZENTRY	AUC <sub>12</sub> (ng.h/mL)	C <sub>avg</sub> (ng/mL)	C <sub>max</sub> (ng/mL)	C <sub>min</sub> (ng/mL)	
10 kg to <20 kg	50 mg twice daily	2,349	196	324	78	
20 kg to <30 kg	75 mg twice daily	3,020	252	394	118	
30 kg to <40 kg	100 mg twice daily	3,229	269	430	126	
≥40 kg	150 mg twice daily	4,044	337	563	152	

<sup>&</sup>lt;sup>a</sup> The covariate distribution of the study population of 85 subjects on CYP3A-inhibitorcontaining regimens was randomly sampled with replacement to obtain 1,000 subjects. Shown in the table are model-predicted steady-state PK parameters for the 1,000 subjects in the simulation dataset.

Table 13. Maraviroc Pharmacokinetic Parameters in Treatment-Experienced Pediatric Patients Receiving SELZENTRY with Noninteracting Concomitant Medications<sup>a</sup>

		Maraviroc Pharmacokinetic Parameter				
			Geometric	Mean		
	Dose of	AUC <sub>12</sub> Cavg Cmax Cmin				
Weight (n)	SELZENTRY	(ng.h/mL)	(ng/mL)	(ng/mL)	(ng/mL)	
<30 kg	Insufficient data					
$\geq 30 \text{ kg } (n = 5)^b$	300 mg	1,998	167	413	50.6	
	twice daily					

<sup>591</sup> <sup>a</sup> Noninteracting concomitant medications include all medications that are not potent CYP3A 592 inhibitors or inducers.

594 Geriatric Patients: Pharmacokinetics of maraviroc have not been fully evaluated in the elderly

595 (aged 65 years and older). Based on population pharmacokinetic analyses, age did not have a

596 clinically relevant effect on maraviroc exposure in subjects up to age 65 years [see Use in

597 *Specific Populations* (8.5)].

598 Race and Gender: Based on population pharmacokinetics and 2 clinical CYP3A5 genotype

599 analyses for race, no dosage adjustment is recommended based on race or gender.

### **Drug Interaction Studies**

601 Effect of Concomitant Drugs on the Pharmacokinetics of Maraviroc: Maraviroc is a substrate of

CYP3A and P-gp and hence its pharmacokinetics are likely to be modulated by inhibitors and 602

<sup>593</sup> <sup>b</sup> Nine observations from 5 subjects.

603 inducers of these enzymes/transporters. The CYP3A/P-gp inhibitors ketoconazole, boceprevir, 604 lopinavir/ritonavir, ritonavir, darunavir/ritonavir, saquinavir/ritonavir, and atazanavir ± ritonavir 605 all increased the C<sub>max</sub> and AUC of maraviroc (Table 14). The CYP3A and/or P-gp inducers rifampin, etravirine, and efavirenz decreased the C<sub>max</sub> and AUC of maraviroc (Table 14). While 606 607 not studied, potent CYP3A and/or P-gp inducers carbamazepine, phenobarbital, and phenytoin 608 are expected to decrease maraviroc concentrations. Based on in vitro study results, maraviroc is 609 also a substrate of OATP1B1 and MRP2; its pharmacokinetics may be modulated by inhibitors of 610 these transporters.

Tipranavir/ritonavir (net CYP3A inhibitor/P-gp inducer) did not affect the steady-state pharmacokinetics of maraviroc (Table 14). Cotrimoxazole and tenofovir did not affect the pharmacokinetics of maraviroc.

611

612

613

614

Table 14. Effect of Coadministered Agents on the Pharmacokinetics of Maraviroc

Coadministered Drug		Dose of	Ratio (90% CI) of Maraviroc Pharmacokinetic Parameters with/without Coadministered Drug (No Effect = 1.00)			
and Dose	n	SELZENTRY	$\mathbf{C}_{\mathbf{min}}$	AUCtau	C <sub>max</sub>	
CYP3A and/or P-gp Inhib	itors					
Ketoconazole	12	100 mg b.i.d.	3.75	5.00	3.38	
400 mg q.d.			(3.01, 4.69)	(3.98, 6.29)	(2.38, 4.78)	
Ritonavir	8	100 mg b.i.d.	4.55	2.61	1.28	
100 mg b.i.d.			(3.37, 6.13)	(1.92, 3.56)	(0.79, 2.09) 4.78	
Saquinavir (soft gel	11	100 mg b.i.d.	11.3	9.77	4.78	
capsules) /ritonavir			(8.96, 14.1)	(7.87, 12.14)	(3.41, 6.71)	
1,000 mg/100 mg b.i.d.						
Lopinavir/ritonavir	11	300 mg b.i.d.	9.24	3.95	1.97	
400 mg/100 mg b.i.d.			(7.98, 10.7)	(3.43, 4.56)	(1.66, 2.34)	
Atazanavir	12	300 mg b.i.d.	4.19	3.57	2.09	
400 mg q.d.		-	(3.65, 4.80)	(3.30, 3.87)	(1.72, 2.55)	
Atazanavir/ritonavir	12	300 mg b.i.d.	6.67	4.88	2.67	
300 mg/100 mg q.d.			(5.78, 7.70)	(4.40, 5.41)	(2.32, 3.08)	
Darunavir/ritonavir	12	150 mg b.i.d.	8.00	4.05	2.29	
600 mg/100 mg b.i.d.			(6.35, 10.1)	(2.94, 5.59)	(1.46, 3.59)	
Boceprevir	14	150 mg b.i.d.	2.78	3.02	3.33	
800 mg t.i.d.			(2.40, 3.23)	(2.53, 3.59)	(2.54, 4.36)	
Elvitegravir/ritonavir	11	150 mg b.i.d.	4.23	2.86	2.15	
150 mg/100 mg q.d.			(3.47, 5.16)	(2.33, 3.51)	(1.71, 2.69)	
CYP3A and/or P-gp Induc	ers					
Efavirenz	12	100 mg b.i.d.	0.55	0.55	0.49	
600 mg q.d.			(0.43, 0.72)	(0.49, 0.62)	(0.38, 0.63)	
Efavirenz	12	200 mg b.i.d.	1.09	1.15	1.16	
600 mg q.d.		(+ efavirenz):	(0.89, 1.35)	(0.98, 1.35)	(0.87, 1.55)	
		100 mg b.i.d.				
		(alone)				
Rifampicin	12	100 mg b.i.d.	0.22	0.37	0.34	
600 mg q.d.			(0.17, 0.28)	(0.33, 0.41)	(0.26, 0.43)	

Rifampicin	12	200 mg b.i.d.	0.66	1.04	0.97
600 mg q.d.		(+ rifampicin):	(0.54, 0.82)	(0.89, 1.22)	(0.72, 1.29)
		100 mg b.i.d.		, , ,	
		(alone)			
Etravirine	14	300 mg b.i.d.	0.61	0.47	0.40
200 mg b.i.d.			(0.53, 0.71)	(0.38, 0.58)	(0.28, 0.57)
Nevirapine <sup>a</sup>	8	300 mg single	_	1.01	1.54
200 mg b.i.d.		dose		(0.65, 1.55)	(0.94, 2.51)
(+ lamivudine 150 mg b.i.d.,					
tenofovir 300 mg q.d.)					
CYP3A and/or P-gp Inhibitors and Inducers					
Lopinavir/ritonavir +	11	300 mg b.i.d.	6.29	2.53	1.25
efavirenz			(4.72, 8.39)	(2.24, 2.87)	(1.01, 1.55)
400 mg/100 mg b.i.d. +					
600 mg q.d.					
Saquinavir(soft gel	11	100 mg b.i.d.	8.42	5.00	2.26
capsules) /ritonavir +			(6.46, 10.97)	(4.26, 5.87)	(1.64, 3.11)
efavirenz					
1,000 mg/100 mg b.i.d. +					
600 mg q.d.					
Darunavir/ritonavir +	10	150 mg b.i.d.	5.27	3.10	1.77
etravirine			(4.51, 6.15)	(2.57, 3.74)	(1.20, 2.60)
600 mg/100 mg b.i.d. +					
200 mg b.i.d.					
Fosamprenavir/ritonavir	14	300 mg b.i.d.	4.74	2.49	1.52
700 mg/100 mg b.i.d.			(4.03, 5.57)	(2.19, 2.82)	(1.27, 1.82)
Fosamprenavir/ritonavir	14	300 mg q.d.	1.80	2.26	1.45
1,400 mg/100 mg q.d.			(1.53, 2.13)	(1.99, 2.58)	(1.20, 1.74)
Tipranavir/ritonavir	12	150 mg b.i.d.	1.80	1.02	0.86
500 mg/200 mg b.i.d.			(1.55, 2.09)	(0.85, 1.23)	(0.61, 1.21)
Other	ı	1	T		
Raltegravir	17	300 mg b.i.d.	0.90	0.86	0.79
400 mg b.i.d.			(0.85, 0.96)	(0.80, 0.92)	(0.67, 0.94)

<sup>615 &</sup>lt;sup>a</sup> Compared with historical data.

Effect of Maraviroc on the Pharmacokinetics of Concomitant Drugs: Maraviroc is unlikely to inhibit the metabolism of coadministered drugs metabolized by the following cytochrome P enzymes (CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, and CYP3A) or to inhibit the uptake of OATP1B1 or the export of MRP2 because maraviroc did not inhibit activity of those enzymes or transporters at clinically relevant concentrations in vitro. Maraviroc does not induce CYP1A2 in vitro. Additionally, in vitro studies have shown that maraviroc is not a substrate for, and does not inhibit, any of the major renal uptake inhibitors (organic anion transporter [OAT]1, OAT3, organic cation transporter [OCT]2, novel organic cation transporter [OCTN]1, and OCTN2) at clinically relevant concentrations.

- In vitro results suggest that maraviroc could inhibit P-gp in the gut. However, maraviroc did not
- significantly affect the pharmacokinetics of digoxin in vivo, indicating maraviroc may not
- significantly inhibit or induce P-gp clinically.
- Drug interaction trials were performed with maraviroc and other drugs likely to be
- 629 coadministered or commonly used as probes for pharmacokinetic interactions (Table 14).
- 630 Coadministration of fosamprenavir 700 mg/ritonavir 100 mg twice daily and maraviroc 300 mg
- twice daily decreased the C<sub>min</sub> and AUC of amprenavir by 36% and 35%, respectively.
- 632 Coadministration of fosamprenavir 1,400 mg/ritonavir 100 mg once daily and maraviroc 300 mg
- once daily decreased the C<sub>min</sub> and AUC by 15% and 30%, respectively. No dosage adjustment is
- 634 necessary when SELZENTRY is dosed 150 mg twice daily in combination with
- 635 fosamprenavir/ritonavir dosed once or twice daily. Fosamprenavir should be given with ritonavir
- when coadministered with SELZENTRY.
- Maraviroc had no significant effect on the pharmacokinetics of elvitegravir, boceprevir,
- 27% and zidovudine, or lamivudine. Maraviroc decreased the C<sub>min</sub> and AUC of raltegravir by 27% and
- 639 37%, respectively, which is not clinically significant. Maraviroc had no clinically relevant effect
- on the pharmacokinetics of midazolam, the oral contraceptives ethinylestradiol and
- 641 levonorgestrel, no effect on the urinary 6β-hydroxycortisol/cortisol ratio, suggesting no induction
- of CYP3A in vivo. Maraviroc had no effect on the debrisoquine metabolic ratio (MR) at 300 mg
- twice daily or less in vivo and did not cause inhibition of CYP2D6 in vitro until concentrations
- greater than 100 microM. However, there was 234% increase in debrisoquine MR on treatment
- compared with baseline at 600 mg once daily, suggesting potential inhibition of CYP2D6 at
- 646 higher doses.

### 647 **12.4 Microbiology**

- Mechanism of Action
- Maraviroc is a member of a therapeutic class called CCR5 co-receptor antagonists. Maraviroc
- selectively binds to the human chemokine receptor CCR5 present on the cell membrane,
- preventing the interaction of HIV-1 gp120 and CCR5 necessary for CCR5-tropic HIV-1 to enter
- cells. CXCR4-tropic and dual-tropic HIV-1 entry is not inhibited by maraviroc.
- Antiviral Activity in Cell Culture
- Maraviroc inhibits the replication of CCR5-tropic laboratory strains and primary isolates of
- 655 HIV-1 in models of acute peripheral blood leukocyte infection. The mean EC<sub>50</sub> value (50%
- effective concentration) for maraviroc against HIV-1 group M isolates (subtypes A to J and
- 657 circulating recombinant form AE) and group O isolates ranged from 0.1 to 4.5 nM (0.05 to
- 658 2.3 ng per mL) in cell culture.
- When used with other antiretroviral agents in cell culture, the combination of maraviroc was not
- antagonistic with non-nucleoside reverse transcriptase inhibitors (NNRTIs: delavirdine,
- efavirenz, and nevirapine), NRTIs (abacavir, didanosine, emtricitabine, lamivudine, stavudine,

- tenofovir, zalcitabine, and zidovudine), or protease inhibitors (PIs: amprenavir, atazanavir,
- darunavir, indinavir, lopinavir, nelfinavir, ritonavir, saquinavir, and tipranavir). Maraviroc was
- not antagonistic with the HIV-1 gp41 fusion inhibitor enfuvirtide. Maraviroc was not active
- against CXCR4-tropic and dual-tropic viruses (EC<sub>50</sub> value greater than 10 microM). The
- antiviral activity of maraviroc against HIV-2 has not been evaluated.
- 667 Resistance in Cell Culture: HIV-1 variants with reduced susceptibility to maraviroc have been
- selected in cell culture following serial passage of 2 CCR5-tropic viruses (CCl/85 and RU570).
- The maraviroc-resistant viruses remained CCR5-tropic with no evidence of a change from a
- 670 CCR5-tropic virus to a CXCR4-using virus. Two amino acid residue substitutions in the V3-loop
- region of the HIV-1 envelope glycoprotein (gp160), A316T, and I323V (HXB2 numbering),
- were shown to be necessary for the maraviroc-resistant phenotype in the HIV-1 isolate CCl/85.
- In the RU570 isolate a 3-amino acid residue deletion in the V3 loop, ΔQAI (HXB2 positions 315
- 674 to 317), was associated with maraviroc resistance. The relevance of the specific gp120
- substitutions observed in maraviroc-resistant isolates selected in cell culture to clinical maraviroc
- 676 resistance is not known. Maraviroc-resistant viruses were characterized phenotypically by
- 677 concentration-response curves that did not reach 100% inhibition in phenotypic drug assays,
- rather than increases in  $EC_{50}$  values.
- 679 Cross-Resistance in Cell Culture: Maraviroc had antiviral activity against HIV-1 clinical isolates
- resistant to NNRTIs, NRTIs, PIs, and the gp41 fusion inhibitor enfuvirtide in cell culture (EC<sub>50</sub>
- values ranged from 0.7 to 8.9 nM [0.36 to 4.57 ng per mL]). Maraviroc-resistant viruses that
- 682 emerged in cell culture remained susceptible to enfuvirtide and the protease inhibitor saquinavir.
- 683 Clinical Resistance: Virologic failure on maraviroc can result from genotypic and phenotypic
- resistance to maraviroc, through outgrowth of undetected CXCR4-using virus present before
- 685 maraviroc treatment (see *Tropism* below), through resistance to background therapy drugs (Table
- 686 15), or due to low exposure to maraviroc [see Clinical Pharmacology (12.2)].
- 687 Antiretroviral Treatment-Experienced Adult Subjects (Trials A4001027 and A4001028): Week
- 48 data from treatment-experienced subjects failing maraviroc-containing regimens with
- 689 CCR5-tropic virus (n = 58) have identified 22 viruses that had decreased susceptibility to
- 690 maraviroc characterized in phenotypic drug assays by concentration-response curves that did not
- reach 100% inhibition. Additionally, CCR5-tropic virus from 2 of these treatment-failure
- subjects had greater than or equal to 3-fold shifts in EC<sub>50</sub> values for maraviroc at the time of
- 693 failure.
- Fifteen of these viruses were sequenced in the gp120 encoding region and multiple amino acid
- substitutions with unique patterns in the heterogeneous V3 loop region were detected. Changes at
- either amino acid position 308 or 323 (HXB2 numbering) were seen in the V3 loop in 7 of the
- subjects with decreased maraviroc susceptibility. Substitutions outside the V3 loop of gp120 may
- also contribute to reduced susceptibility to maraviroc.

699 Antiretroviral Treatment-Naive Adult Subjects (Trial A4001026): Treatment-naive subjects 700 receiving SELZENTRY had more virologic failures and more treatment-emergent resistance to 701 the background regimen drugs compared with those receiving efavirenz (Table 15).

Table 15. Development of Resistance to Maraviroc or Efavirenz and Background Drugs in Antiretroviral Treatment-Naive Trial A4001026 for Patients with Only CCR5-Tropic Virus at Screening Using Enhanced Sensitivity TROFILE Assay

Maraviroc Efavirenz Total N in dataset (as-treated) 273 241 Total virologic failures (as-treated) 85 (31%) 56 (23%) 43 Evaluable virologic failures with post baseline 73 genotypic and phenotypic data Lamivudine resistance 39 (53%) 13 (30%) Zidovudine resistance 2 (3%) 0 Efavirenz resistance 23 (53%) Phenotypic resistance to maraviroc<sup>a</sup> 19 (26%)

707 In an as-treated analysis of treatment-naive subjects at 96 weeks, 32 subjects failed a 708 maraviroc-containing regimen with CCR5-tropic virus and had a tropism result at failure; 7 of 709 these subjects had evidence of maraviroc phenotypic resistance defined as 710 concentration-response curves that did not reach 95% inhibition. One additional subject had a 711 greater than or equal to 3-fold shift in the EC<sub>50</sub> value for maraviroc at the time of failure. A 712 clonal analysis of the V3 loop amino acid envelope sequences was performed from 6 of the 713 7 subjects. Changes in V3 loop amino acid sequence differed between each of these different 714 subjects, even for those infected with the same virus clade, suggesting that there are multiple 715 diverse pathways to maraviroc resistance. The subjects who failed with CCR5-tropic virus and 716 without a detectable maraviroc shift in susceptibility were not evaluated for genotypic resistance.

717 Of the 32 maraviroc virologic failures failing with CCR5-tropic virus, 20 (63%) also had 718

genotypic and/or phenotypic resistance to background drugs in the regimen (lamivudine,

719 zidovudine).

702

703

704

720 Tropism: In both treatment-experienced and treatment-naive subjects, detection of CXCR4-using

721 virus prior to initiation of therapy has been associated with a reduced virologic response to

722 maraviroc.

723 Antiretroviral Treatment-Experienced Subjects (Trials A4001027 and A4001028): In the

724 majority of cases, treatment failure on maraviroc was associated with detection of CXCR4-using

725 virus (i.e., CXCR4- or dual/mixed-tropic) which was not detected by the tropism assay prior to

726 treatment. CXCR4-using virus was detected at failure in approximately 55% of subjects who

<sup>705</sup> <sup>a</sup> Includes subjects failing with CXCR4- or dual/mixed-tropism because these viruses are not 706 intrinsically susceptible to maraviroc.

- failed treatment on maraviroc by Week 48, as compared with 9% of subjects who experienced
- treatment failure in the placebo arm. To investigate the likely origin of the on-treatment
- 729 CXCR4-using virus, a detailed clonal analysis was conducted on virus from 20 representative
- subjects (16 subjects from the maraviroc arms and 4 subjects from the placebo arm) in whom
- 731 CXCR4-using virus was detected at treatment failure. From analysis of amino acid sequence
- differences and phylogenetic data, it was determined that CXCR4-using virus in these subjects
- emerged from a low level of pre-existing CXCR4-using virus not detected by the tropism assay
- (which is population-based) prior to treatment rather than from a co-receptor switch from
- 735 CCR5-tropic virus to CXCR4-using virus resulting from mutation in the virus.
- 736 Detection of CXCR4-using virus prior to initiation of therapy has been associated with a reduced
- virological response to maraviroc. Furthermore, subjects failing twice-daily maraviroc at Week
- 48 with CXCR4-using virus had a lower median increase in CD4+ cell counts from baseline
- 739 (+41 cells per mm<sup>3</sup>) than those subjects failing with CCR5-tropic virus (+162 cells per mm<sup>3</sup>).
- 740 The median increase in CD4+ cell count in subjects failing in the placebo arm was +7 cells per
- 741  $\text{mm}^3$ .
- 742 Antiretroviral Treatment-Naive Subjects (Trial A4001026): In a 96-week trial of antiretroviral
- 743 treatment-naive subjects, 14% (12 of 85) who had only CCR5-tropic virus at screening with an
- enhanced sensitivity tropism assay (TROFILE) and failed therapy on maraviroc had
- 745 CXCR4-using virus at the time of treatment failure. A detailed clonal analysis was conducted in
- 746 2 previously antiretroviral treatment-naive subjects enrolled in a Phase 2a monotherapy trial who
- had CXCR4-using virus detected after 10 days' treatment with maraviroc. Consistent with the
- detailed clonal analysis conducted in treatment-experienced subjects, the CXCR4-using variants
- appear to emerge from outgrowth of a pre-existing undetected CXCR4-using virus. Screening
- with an enhanced sensitivity tropism assay reduced the number of maraviroc virologic failures
- with CXCR4- or dual/mixed-tropic virus at failure to 12 compared with 24 when screening with
- 752 the original tropism assay. All but one (11 of 12; 92%) of the maraviroc failures failing with
- 753 CXCR4- or dual/mixed-tropic virus also had genotypic and phenotypic resistance to the
- background drug lamivudine at failure and 33% (4 of 12) developed zidovudine-associated
- 755 resistance substitutions.
- 756 Subjects who had only CCR5-tropic virus at baseline and failed maraviroc therapy with
- CXCR4-using virus had a median increase in CD4+ cell counts from baseline of +113 cells per
- 758 mm<sup>3</sup> while those subjects failing with CCR5-tropic virus had an increase of +135 cells per mm<sup>3</sup>.
- 759 The median increase in CD4+ cell count in subjects failing in the efavirenz arm was +95 cells
- 760 per  $mm^3$ .
- 761 Antiretroviral Treatment-Experienced Pediatric Subjects (Trial A4001031): In the Week 48
- analysis of Trial A4001031 (n = 103), the mechanisms of resistance to maraviroc observed in the
- treatment-experienced pediatric population were similar to those observed in adult populations:
- reasons for virologic failure included failing with CXCR4- or dual/mixed-tropic virus, evidence

- of reduced maraviroc susceptibility as measured by a decrease in maximal percentage inhibition
- 766 (MPI), and emergence of resistance to background drug in the regimen.

#### 767 13 NONCLINICAL TOXICOLOGY

# 768 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

- 769 <u>Carcinogenesis</u>
- Long-term oral carcinogenicity studies of maraviroc were carried out in rasH2 transgenic mice
- 771 (6 months) and in rats for up to 96 weeks (females) and 104 weeks (males). No drug-related
- increases in tumor incidence were found in mice at 1,500 mg per kg per day and in male and
- female rats at 900 mg per kg per day. The highest exposures in rats were approximately 11 times
- those observed in humans at the therapeutic dose of 300 mg twice daily for the treatment of
- 775 HIV-1 infection.
- 776 <u>Mutagenesis</u>
- Maraviroc was not genotoxic in the reverse mutation bacterial test (Ames test in Salmonella and
- E. coli), a chromosome aberration test in human lymphocytes, and mouse bone marrow
- 779 micronucleus test.
- 780 <u>Impairment of Fertility</u>
- Maraviroc did not impair mating or fertility of male or female rats and did not affect sperm of
- treated male rats at approximately 20-fold higher exposures (AUC) than in humans given the
- 783 recommended 300-mg twice-daily dose.

#### 784 14 CLINICAL STUDIES

#### 785 **14.1** Clinical Studies in Adult Subjects

- The clinical efficacy and safety of SELZENTRY are derived from analyses of data from 3 trials
- in adult subjects infected with CCR5-tropic HIV-1: Trials A4001027 and A4001028 in
- antiretroviral treatment-experienced adult subjects and Trial A4001026 in treatment-naive
- subjects. These trials were supported by a 48-week trial in antiretroviral treatment-experienced
- adult subjects infected with dual/mixed-tropic HIV-1, Trial A4001029.

#### 791 Trials in CCR5-Tropic, Treatment-Experienced Subjects

- 792 Trials A4001027 and A4001028 were double-blind, randomized, placebo-controlled, multicenter
- trials in subjects infected with CCR5-tropic HIV-1. Subjects were required to have an HIV-1
- RNA greater than 5,000 copies per mL despite at least 6 months of prior therapy with at least
- 1 agent from 3 of the 4 antiretroviral drug classes (greater than or equal to 1 NRTI, greater than
- or equal to 1 NNRTI, greater than or equal to 2 PIs, and/or enfuvirtide) or documented resistance
- to at least 1 member of each class. All subjects received an optimized background regimen
- 798 consisting of 3 to 6 antiretroviral agents (excluding low-dose ritonavir) selected on the basis of

the subject's prior treatment history and baseline genotypic and phenotypic viral resistance measurements. In addition to the optimized background regimen, subjects were then randomized in a 2:2:1 ratio to SELZENTRY 300 mg once daily, SELZENTRY 300 mg twice daily, or placebo. Doses were adjusted based on background therapy as described in *Dosage and Administration* (2), Table 1.

In the pooled analysis for Trials A4001027 and A4001028, the demographics and baseline

In the pooled analysis for Trials A4001027 and A4001028, the demographics and baseline characteristics of the treatment groups were comparable (Table 16). Of the 1,043 subjects with a CCR5-tropism result at screening, 7.6% had a dual/mixed-tropism result at the baseline visit 4 to 6 weeks later. This illustrates the background change from CCR5- to dual/mixed-tropism result over time in this treatment-experienced population, prior to a change in antiretroviral regimen or administration of a CCR5 co-receptor antagonist.

Table 16. Demographic and Baseline Characteristics of Subjects in Trials A4001027 and A4001028

	SELZENTRY	
	Twice Daily	Placebo
	(n = 426)	(n = 209)
Age (years)		
Mean (range)	46.3 (21-73)	45.7 (29-72)
Sex:		
Male	382 (89.7%)	185 (88.5%)
Female	44 (10.3%)	24 (11.5%)
Race:		
White	363 (85.2%)	178 (85.2%)
Black	51 (12.0%)	26 (12.4%)
Other	12 (2.8%)	5 (2.4%)
Region:		
U.S.	276 (64.8%)	135 (64.6%)
Non-U.S.	150 (35.2%)	74 (35.4%)
Subjects with previous enfuvirtide use	142 (33.3%)	62 (29.7%)
Subjects with enfuvirtide as part of OBT	182 (42.7%)	91 (43.5%)
Baseline plasma HIV-1 RNA (log <sub>10</sub> copies/mL)		
Mean (range)	4.85 (2.96-6.88)	4.86 (3.46-7.07)
Subjects with screening viral load ≥100,000 copies/mL	179 (42.0%)	84 (40.2%)
Baseline CD4+ cell count (cells/mm <sup>3</sup> )		
Median (range)	167 (2-820)	171 (1-675)
Subjects with baseline CD4+ cell count ≤200 cells/mm <sup>3</sup> )	250 (58.7%)	118 (56.5%)
Subjects with Overall Susceptibility Score (OSS): <sup>a</sup>		
0	57 (13.4%)	35 (16.7%)
1	136 (31.9%)	44 (21.1%)

2	104 (24.4%)	59 (28.2%)
≥3	125 (29.3%)	66 (31.6%)
Subjects with enfuvirtide resistance substitutions	90 (21.2%)	45 (21.5%)
Median number of resistance-associated: <sup>b</sup>		
PI substitutions	10	10
NNRTI substitutions	1	1
NRTI substitutions	6	6

a OSS - Sum of active drugs in OBT based on combined information from genotypic and phenotypic testing.

Table 17. Outcomes of Randomized Treatment at Week 48 in Trials A4001027 and A4001028

	SELZENTRY Twice Daily	Placebo	Mean
Outcome	(n = 426)	(n = 209)	Difference
Mean change from Baseline to Week 48 in	-1.84	-0.78	-1.05
HIV-1 RNA (log <sub>10</sub> copies/mL)			
<400 copies/mL at Week 48	239 (56%)	47 (22%)	34%
<50 copies/mL at Week 48	194 (46%)	35 (17%)	29%
Discontinuations:			
Insufficient clinical response	97 (23%)	113 (54%)	_
Adverse events	19 (4%)	11 (5%)	_
Other	27 (6%)	18 (9%)	_
Subjects with treatment-emergent CDC	22 (5%)	16 (8%)	_
Category C events			
Deaths (during trial or within 28 days of	9 (2%) <sup>a</sup>	1 (0.5%)	_
last dose)			

<sup>&</sup>lt;sup>a</sup> One additional subject died while receiving open-label therapy with SELZENTRY subsequent to discontinuing double-blind placebo due to insufficient response.

After 48 weeks of therapy, the proportions of subjects with HIV-1 RNA less than 400 copies per mL receiving SELZENTRY compared with placebo were 56% and 22%, respectively. The mean changes in plasma HIV-1 RNA from baseline to Week 48 were –1.84 log<sub>10</sub> copies per mL for subjects receiving SELZENTRY + OBT compared with –0.78 log<sub>10</sub> copies per mL for subjects receiving OBT only. The mean increase in CD4+ cell count was higher on SELZENTRY twice daily + OBT (124 cells per mm<sup>3</sup>) than on placebo + OBT (60 cells per mm<sup>3</sup>).

Trial in Dual/Mixed-Tropic, Treatment-Experienced Subjects

<sup>814</sup> b Resistance substitutions based on IAS guidelines.<sup>1</sup>

The Week 48 results for the pooled Trials A4001027 and A4001028 are shown in Table 17.

- 827 Trial A4001029 was an exploratory, randomized, double-blind, multicenter trial to determine the 828 safety and efficacy of SELZENTRY in subjects infected with dual/mixed co-receptor tropic 829 HIV-1. The inclusion/exclusion criteria were similar to those for Trials A4001027 and A4001028 above and the subjects were randomized in a 1:1:1 ratio to SELZENTRY once daily, 830 831 SELZENTRY twice daily, or placebo. No increased risk of infection or HIV-1 disease 832 progression was observed in the subjects who received SELZENTRY. Use of SELZENTRY was 833 not associated with a significant decrease in HIV-1 RNA compared with placebo in these 834 subjects and no adverse effect on CD4+ cell count was noted. 835 Trial in Treatment-Naive Subjects 836 Trial A4001026 was a randomized, double-blind, multicenter trial in subjects infected with 837 CCR5-tropic HIV-1 classified by the original TROFILE tropism assay. Subjects were required to 838 have plasma HIV-1 RNA greater than or equal to 2,000 copies per mL and could not have: 1) 839 previously received any antiretroviral therapy for greater than 14 days, 2) an active or recent 840 opportunistic infection or a suspected primary HIV-1 infection, or 3) phenotypic or genotypic resistance to zidovudine, lamivudine, or efavirenz. Subjects were randomized in a 1:1:1 ratio to 841 842 SELZENTRY 300 mg once daily, SELZENTRY 300 mg twice daily, or efavirenz 600 mg once
- pre-specified criteria for demonstrating non-inferiority and was discontinued.

  The demographic and baseline characteristics of the maraviroc and efavirenz treatment groups were comparable (Table 18). Subjects were stratified by screening HIV-1 RNA levels and by geographic region. The median CD4+ cell counts and mean HIV-1 RNA at baseline were similar for both treatment groups.

SELZENTRY are based on the comparison of SELZENTRY twice daily versus efavirenz. In a

pre-planned interim analysis at 16 weeks, SELZENTRY 300 mg once daily failed to meet the

daily, each in combination with lamivudine/zidovudine. The efficacy and safety of

843

844

845

851

Table 18. Demographic and Baseline Characteristics of Subjects in Trial A4001026

	SELZENTRY	Efavirenz
	300 mg Twice Daily +	600 mg Once Daily +
	Lamivudine/Zidovudine	Lamivudine/Zidovudine
	(n = 360)	(n = 361)
Age (years):		
Mean	36.7	37.4
Range	20-69	18-77
Female, n%	104 (29)	102 (28)
Race, n%:		
White	204 (57)	198 (55)
Black	123 (34)	133 (37)
Asian	6 (2)	5 (1)
Other	27 (8)	25 (7)

Median (range) CD4+ cell count	241 (5-1,422)	254 (8-1,053)
(cells/microL)		
Median (range) HIV-1 RNA	4.9 (3-7)	4.9 (3-7)
(log <sub>10</sub> copies/mL)		

The treatment outcomes at 96 weeks for Trial A4001026 are shown in Table 19. Treatment outcomes are based on reanalysis of the screening samples using a more sensitive tropism assay, enhanced sensitivity TROFILE HIV tropism assay, which became available after the Week 48 analysis; approximately 15% of the subjects identified as CCR5-tropic in the original analysis had dual/mixed- or CXCR4-tropic virus. Screening with enhanced sensitivity version of the TROFILE tropism assay reduced the number of maraviroc virologic failures with CXCR4- or dual/mixed-tropic virus at failure to 12 compared with 24 when screening with the original TROFILE HIV tropism assay.

Table 19. Trial Outcome (Snapshot) at Week 96 Using Enhanced Sensitivity Assay<sup>a</sup>

	CDI ZENIDAZ		
	SELZENTRY	Efavirenz	
	300 mg Twice Daily +	600 mg Once Daily +	
	Lamivudine/Zidovudine	Lamivudine/Zidovudine	
	(n = 311)	(n = 303)	
Outcome at Week 96b	n (%)	n (%)	
Virologic Responders:			
(HIV-1 RNA <400 copies/mL)	199 (64)	195 (64)	
Virologic Failure:			
Non-sustained HIV-1 RNA	39 (13)	22 (7)	
suppression			
HIV-1 RNA never suppressed	9 (3)	1 (<1)	
Virologic Responders:			
(HIV-1 RNA <50 copies/mL)	183 (59)	190 (63)	
Virologic Failure:			
Non-sustained HIV-1 RNA	43 (14)	25 (8)	
suppression	- ( )	- (-)	
HIV-1 RNA never suppressed	21 (7)	3 (1)	
Discontinuations due to:			
Adverse events	19 (6)	47 (16)	
Death	2(1)	2(1)	
Other <sup>c</sup>	43 (14)	36 (12)	

<sup>a</sup> The total number of subjects (311, 303) in Table 19 represents the subjects who had a CCR5-tropic virus in the reanalysis of screening samples using the more sensitive tropism assay. This reanalysis reclassified approximately 15% of subjects shown in Table 18 as having dual/mixed- or CXCR4-tropic virus. These numbers are different than those presented in Table 18 because the numbers in Table 18 reflect the subjects with CCR5-tropic virus according to

- the original tropism assay.
- b Week 48 results: Virologic responders (less than 400): 228 of 311 (73%) in SELZENTRY, 219
- 868 of 303 (72%) in efavirenz;
- 869 Virologic responders (less than 50): 213 of 311 (69%) in SELZENTRY, 207 of 303 (68%) in
- 870 efavirenz
- 871 ° Other reasons for discontinuation include lost to follow-up, withdrawn, protocol violation, and
- other.
- The median increase from baseline in CD4+ cell counts at Week 96 was 184 cells per mm<sup>3</sup> for
- the arm receiving SELZENTRY compared with 155 cells per mm<sup>3</sup> for the efavirenz arm.

# 875 14.2 Clinical Studies in Pediatric Subjects

- 876 Trial in CCR5-Tropic, Treatment-Experienced Subjects
- 877 Trial A4001031 is an open-label, multicenter trial in pediatric subjects aged 2 to less than
- 878 18 years infected with only CCR5-tropic HIV-1. Subjects were required to have HIV-1 RNA
- greater than 1,000 copies per mL at screening. All subjects (n = 103) received SELZENTRY
- twice daily and OBT. Dosing of SELZENTRY was based on BSA and doses were adjusted
- based on whether the subject was receiving potent CYP3A inhibitors and/or inducers.
- The population was 52% female and 69% black, with mean age of 10 years (range: 2 to
- 17 years). At baseline, mean plasma HIV-1 RNA was 4.4 log<sub>10</sub> copies per mL (range: 2.4 to
- 6.2 log<sub>10</sub> copies per mL), mean CD4+ cell count was 551 cells per mm<sup>3</sup> (range: 1 to
- 1,654 cells per mm<sup>3</sup>), and mean CD4+ percent was 21% (range: 0% to 42%).
- At 48 weeks, 48% of subjects treated with SELZENTRY and OBT achieved plasma HIV-1 RNA
- less than 48 copies per mL and 65% of subjects achieved plasma HIV-1 RNA less than
- 400 copies per mL. The mean CD4+ cell count (percent) increase from baseline to Week 48 was
- 889 247 cells per mm<sup>3</sup> (5%).

#### 890 15 REFERENCES

- 1. IAS-USA Drug Resistance Mutations Figures.
- http://www.iasusa.org/pub/topics/2006/issue3/125.pdf

#### 893 16 HOW SUPPLIED/STORAGE AND HANDLING

- 894 SELZENTRY film-coated tablets are available as follows:
- 895 25-mg, 75-mg, 150-mg, and 300-mg tablets are blue, biconvex, oval, film-coated tablets
- debossed with "MVC 25", "MVC 75", "MVC 150", or "MVC 300", respectively, on one side
- and plain on the other.
- 898 25-mg tablets: Bottle of 120 tablets (NDC 49702-233-08).
- 899 75-mg tablets: Bottle of 120 tablets (NDC 49702-235-08).

- 900 150-mg tablets: Bottle of 60 tablets (NDC 49702-223-18).
- 901 300-mg tablets: Bottle of 60 tablets (NDC 49702-224-18).
- 902 SELZENTRY film-coated tablets should be stored at 20°C to 25°C (68°F to 77°F); excursions
- 903 permitted between 15°C and 30°C (59°F and 86°F) [see USP Controlled Room Temperature].
- 904 SELZENTRY oral solution is a clear, colorless, strawberry-flavored liquid. Each mL of the
- solution contains 20 mg of maraviroc. It is packaged in plastic bottles as follows:
- 906 Bottle of 230 mL (NDC 49702-237-55). Each bottle is packaged with one press-in bottle adapter
- and one 10–mL oral dosing syringe with 0.5–mL gradations. The press-in bottle adapter and oral
- dosing syringe are not made with natural rubber latex. This product does not require
- 909 reconstitution.
- 910 SELZENTRY oral solution should be stored at 20°C to 25°C (68°F to 77°F); excursions
- 911 permitted between 15°C and 30°C (59°F and 86°F) [see USP Controlled Room Temperature].
- 912 Discard any unused oral solution 60 days after first opening the bottle.

### 913 17 PATIENT COUNSELING INFORMATION

- Advise the patient to read the FDA-approved patient labeling (Medication Guide and Instructions
- 915 for Use).
- 916 Hepatotoxicity
- Inform patients that hepatotoxicity, including life-threatening cases, has been reported with
- 918 SELZENTRY; therefore, it is important to inform the healthcare professional if patients have
- 919 underlying hepatitis B or C or elevations in liver-associated tests prior to treatment. Inform
- patients to stop SELZENTRY and seek medical evaluation immediately if they develop signs or
- 921 symptoms of hepatitis or allergic reaction following use of SELZENTRY. Advise patients that
- laboratory tests for liver enzymes and bilirubin will be ordered prior to starting SELZENTRY, at
- other times during treatment, and if they develop severe rash or signs and symptoms of hepatitis
- or an allergic reaction on treatment [see Dosage and Administration (2.1), Warnings and
- 925 *Precautions* (5.1, 5.2)].
- 926 Cardiovascular Events
- When administering SELZENTRY in patients with cardiovascular comorbidities, a history of
- 928 postural hypotension or receiving concomitant medication known to lower blood pressure, advise
- patients that they may be at increased risk for cardiovascular events. Advise patients to avoid
- 930 driving or operating machinery if they experience dizziness while taking SELZENTRY [see
- 931 *Warnings and Precautions* (5.3)].
- 932 Drug Interactions
- Advise patients to inform their healthcare provider of concomitant HIV medications as dosage of
- 934 SELZENTRY may be modified depending on other HIV medications taken with SELZENTRY.

<ul><li>935</li><li>936</li><li>937</li></ul>	Advise patients that coadministration of SELZENTRY with St. John's wort is not recommended as it can lead to loss of virologic response and possible resistance to SELZENTRY [see Dosage and Administration (2.2), Drug Interactions (7.1)].
938	Missed Dosage
939 940 941 942 943	Inform patients that it is important to take SELZENTRY in combination with other antiretroviral medications on a regular dosing schedule with or without food. Advise patients to avoid missing doses as it can result in development of resistance. Instruct patients that if they miss a dose, to take it as soon as they remember. Advise patients not to double their next dose or take more than the prescribed dose [see Dosage and Administration (2.2)].
944	Pregnancy
945 946 947	Inform patients that there is insufficient data on the safety of SELZENTRY in pregnancy. Inform patients that there is an antiretroviral pregnancy registry that monitors pregnancy outcomes in women exposed to SELZENTRY during pregnancy [see Use in Specific Populations (8.1)].
948	<u>Lactation</u>
949 950	Instruct women with HIV-1 infection not to breastfeed because HIV-1 can be passed to the baby in breast milk [see Use in Specific Populations (8.2)].
951	
952 953	SELZENTRY and COMBIVIR are trademarks owned by or licensed to the ViiV Healthcare group of companies.
954 955 956	TROFILE is a trademark owned by or licensed to Monogram BioSciences, Inc., and is not owned by or licensed to the ViiV Healthcare group of companies. The maker of this brand is not affiliated with and does not endorse the ViiV Healthcare group of companies or its products.
957 958	
959	Manufactured for:
	ViiV
960	Healthcare
961	ViiV Healthcare
962	Research Triangle Park, NC 27709
963	
964	©2018 ViiV Healthcare group of companies or its licensor.
965	SEL:13PI

#### PHARMACIST-DETACH HERE AND GIVE MEDICATION GUIDE TO PATIENT

#### **MEDICATION GUIDE**

SELZENTRY (sell-ZEN-tree) (maraviroc)

tablets

SELZENTRY (sell-ZEN-tree) (maraviroc)

oral solution

What is the most important information I should know about SELZENTRY?

SELZENTRY can cause serious side effects including serious liver problems (liver toxicity). An allergic reaction may happen before liver problems occur. Stop taking SELZENTRY and call your healthcare provider right away if you get any of the following signs or symptoms of liver problems:

- an itchy rash on your body (allergic reaction)
- your skin or the white part of your eyes turns yellow (jaundice)
- dark or "tea-colored" urine

- vomiting
- pain, aching, or tenderness on the right side of your stomach area

Your healthcare provider will do blood tests to check your liver before you begin treatment with SELZENTRY and as needed during treatment, and if you get a severe rash, signs and symptoms of liver problems, or an allergic reaction during treatment with SELZENTRY.

#### What is SELZENTRY?

SELZENTRY is a prescription HIV-1 (Human Immunodeficiency Virus type 1) medicine used with other antiretroviral medicines to treat CCR5-tropic HIV-1 infection in people 2 years of age and older weighing at least 22 lb (10 kg). HIV-1 is the virus that causes AIDS (Acquired Immune Deficiency Syndrome). Use of SELZENTRY is not recommended in people with dual/mixed or CXCR4-tropic HIV-1.

The safety and effectiveness of SELZENTRY has not been established in children younger than 2 years of age.

#### Who should not take SELZENTRY?

## Do not take SELZENTRY if you:

• have severe kidney problems or are on hemodialysis and are also taking certain other medications.

#### What should I tell my healthcare provider before taking SELZENTRY?

Before you take SELZENTRY, tell your healthcare provider about all of your medical conditions, including if you:

- have or have had liver problems including hepatitis B or C virus infection.
- have heart problems.
- have kidney problems.
- have low blood pressure or take medicines to lower blood pressure.
- are pregnant or plan to become pregnant. It is not known if SELZENTRY may harm your unborn baby.
   Pregnancy Registry. There is a pregnancy registry for women who take antiretroviral medicines during pregnancy. The purpose of this registry is to collect information about the health of you and your baby.

Talk to your healthcare provider about how you can take part in this registry.

are breastfeeding or plan to breastfeed. Do not breastfeed if you take SELZENTRY. You should
not breastfeed if you have HIV-1 because of the risk of passing HIV-1 to your baby. Talk to your
healthcare provider about the best way to feed your baby.

Tell your healthcare provider about all the medicines you take, including prescription and over-thecounter medicines, vitamins, and herbal supplements.

Some medicines may interact with SELZENTRY. **Keep a list of your medicines to show your healthcare** provider and pharmacist.

 You can ask your healthcare provider or pharmacist for a list of medicines that interact with SELZENTRY.

**Do not start taking a new medicine without telling your healthcare provider.** Your healthcare provider can tell you if it is safe to take SELZENTRY with other medicines. Your healthcare provider may need to change your dose of SELZENTRY when you take it with certain medicines.

• You should not take SELZENTRY if you also take St. John's wort (Hypericum perforatum).

#### How should I take SELZENTRY?

- Take SELZENTRY exactly as your healthcare provider tells you.
- Do not change your dose or stop taking SELZENTRY without first talking with your healthcare provider.
- If you miss a dose of SELZENTRY, take it as soon as you remember. Do not take 2 doses at the same time. If you are not sure about your dosing, call your healthcare provider.
- Stay under the care of a healthcare provider while taking SELZENTRY.
- Swallow SELZENTRY tablets whole. Do not chew the tablets.
- SELZENTRY may be taken with or without food.
- For children aged 2 years and older and weighing at least 22 lb (10 kg), your healthcare provider will
  prescribe a dose of SELZENTRY based on your child's body weight and other medicines they are
  taking.
- Tell your healthcare provider if your child has trouble swallowing tablets. SELZENTRY comes as tablets or as a liquid (oral solution).
- SELZENTRY oral solution should be given with the supplied press-in bottle adapter and oral dosing syringe. See the Instructions for Use that comes with SELZENTRY oral solution for information about the right way to take a dose.
- Do not run out of SELZENTRY. The virus in your blood may increase and the virus in your blood may become harder to treat. When your supply starts to run low, get more from your healthcare provider or pharmacy.
- If you take too much SELZENTRY, call your healthcare provider or go to the nearest hospital emergency room right away.

#### What are the possible side effects of SELZENTRY?

- SELZENTRY can cause serious side effects including:
- See "What is the most important information I should know about SELZENTRY?"
- Serious skin rash and allergic reactions. Severe and potentially life-threatening skin reactions and allergic reactions have been reported in some patients taking SELZENTRY. If you develop a rash with

any of the following symptoms, stop using SELZENTRY and contact your doctor right away:

- fever
- · generally ill feeling
- muscle aches
- blisters or sores in your mouth
- blisters or peeling of the skin
- redness or swelling of the eyes
- swelling of the mouth or face or lips

- problems breathing
- yellowing of the skin or whites of your eyes
- · dark or tea-colored urine
- pain, aching, or tenderness on the right side below the ribs
- loss of appetite
- nausea/vomiting
- **Heart problems** including heart attack.
- Low blood pressure when standing up (postural hypotension) can cause dizziness or fainting. You should avoid driving or operating heavy machinery if you have dizziness while taking SELZENTRY.
- Changes in your immune system (Immune Reconstitution Syndrome) can happen when you start taking HIV-1 medicines. Your immune system may get stronger and begin to fight infections that have been hidden in your body for a long time. Tell your healthcare provider right away if you develop new symptoms after you start taking SELZENTRY.
- Possible chance of infection or cancer. SELZENTRY affects other immune system cells and therefore may possibly increase your chance for getting other infections or cancer.

The most common side effects of SELZENTRY in adults include colds and cold-like symptoms, cough, fever, rash, bloating and gas, indigestion, constipation, and dizziness.

The most common side effects of SELZENTRY in children include vomiting, abdominal pain, diarrhea, nausea, and dizziness.

Tell your healthcare provider if you have any side effect that bothers you or that does not go away. These are not all the possible side effects of SELZENTRY. For more information, ask your healthcare provider or pharmacist. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

#### How should I store SELZENTRY?

- Store SELZENTRY tablets and oral solution at room temperature between 68°F to 77°F (20°C to 25°C).
- Throw away any unused oral solution 60 days after first opening the bottle.

#### Keep SELZENTRY and all medicines out of the reach of children.

#### **General information about SELZENTRY**

Medicines are sometimes prescribed for purposes other than those mentioned in a Medication Guide. Do not use SELZENTRY for a condition for which it was not prescribed. Do not give SELZENTRY to other people, even if they have the same symptoms that you have. It may harm them.

If you would like more information, talk with your healthcare provider. You can ask your healthcare provider or pharmacist for the information about SELZENTRY that is written for health professionals.

For more information go to www.selzentry.com.

#### What are the ingredients in SELZENTRY?

Active ingredient: maraviroc

Inactive ingredients:

Tablets: Dibasic calcium phosphate (anhydrous), magnesium stearate, microcrystalline cellulose, and sodium starch glycolate. Tablet film-coating contains: FD&C blue #2 aluminum lake, soya lecithin, polyethylene glycol (macrogol 3350), polyvinyl alcohol, talc, and titanium dioxide.

Oral Solution: Citric acid, purified water, sodium benzoate, sodium citrate dihydrate, strawberry flavoring (501440T), and sucralose.

Manufactured for:



ViiV Healthcare

Research Triangle Park, NC 27709

SELZENTRY is a trademark owned by or licensed to the ViiV Healthcare group of companies. ©2018 ViiV Healthcare group of companies or its licensor.

SEL:8MG

This Medication Guide has been approved by the U.S. Food and Drug Administration.

Revised: 07/2018

969970

# INSTRUCTIONS FOR USE SELZENTRY (sell-ZEN-tree) (maraviroc) oral solution

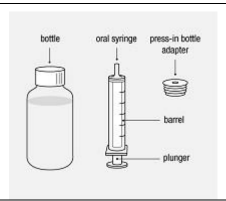
Read this Instructions for Use before you start taking SELZENTRY oral solution and each time you get a refill. There may be new information. This leaflet does not take the place of talking to your healthcare provider about your medical condition or treatment.

#### Important information about measuring SELZENTRY oral solution:

Always use the oral syringe that comes with your SELZENTRY oral solution to measure your prescribed dose. Ask your healthcare provider or pharmacist to show you how to measure your prescribed dose if you are not sure.

# Each carton of SELZENTRY oral solution contains:

- 1 oral dosing syringe
- 1 press-in bottle adapter
- 1 bottle of SELZENTRY oral solution



Before each use: Wash your hands with soap and water and place the items from the carton on a clean flat surface.

# Step 1. Open the bottle of SELZENTRY oral solution.

Open the bottle by pushing down firmly on the childresistant cap and turning it counter-clockwise. **See Figure A.** 

Do not throw away the child-resistant cap.

# Step 2. First time use only: Insert the press-in bottle adapter.

Remove the press-in bottle adapter and oral syringe from the plastic overwrap. With the bottle on a flat surface, push the ribbed end of the press-in bottle adapter all the way into the neck of the bottle while holding the bottle firmly. **See Figure B.** 

Note: Do not remove the press-in bottle adapter from the bottle after it is inserted.

# Step 3. Find your prescribed dose on the oral syringe.

Check the dose in milliliters (mL) as prescribed by your healthcare provider. Find this marking on the oral syringe. **See Figure C.** 

### Step 4. Remove air from oral syringe.

Push the oral syringe plunger to the bottom of the barrel of the syringe (toward its tip) to remove excess air. **See Figure D.** 

Figure A. Opening the bottle



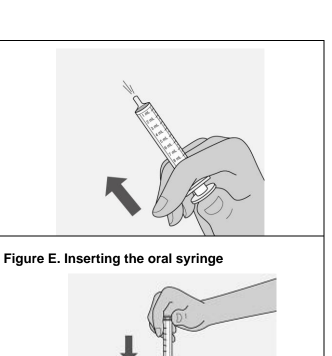
Figure B. Inserting the press-in bottle adapter



Figure C. Find your prescribed dose



Figure D. Removing air from oral syringe.



Step 5. Insert the oral syringe.

Insert the oral syringe into the upright bottle through the opening of the press-in bottle adapter until it is firmly in place. **See Figure E**.

# Step 6. Withdraw the prescribed dose of SELZENTRY from the bottle.

With the oral syringe in place, turn the bottle upside down. Pull back the plunger of the oral syringe until the top of the plunger is even with the markings on the oral syringe for your prescribed dose. **See Figure F.** 

If you see air bubbles in the oral syringe, fully push the plunger in to empty the oral solution back into the bottle. Then withdraw your prescribed dose of oral solution.

### Step 7. Removing the oral syringe.

Turn the bottle upright and place the bottle on a flat surface. Remove the oral syringe from the bottle adapter and bottle by pulling straight up on the oral syringe. **See Figure G.** 

Figure F. Withdrawing the oral solution

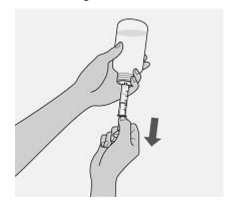
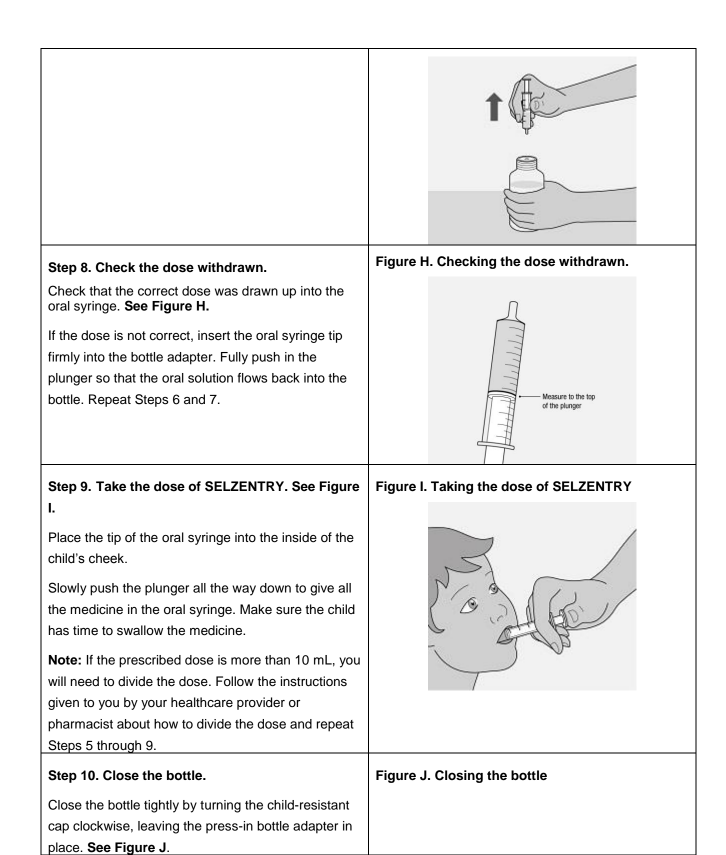
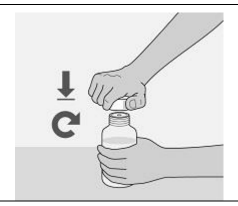


Figure G. Removing the oral syringe





### Step 11. Clean the oral syringe.

Rinse the oral syringe with tap water after each use.

Remove the plunger from the barrel by pulling the plunger and the barrel away from each other. **See Figure K**.

Rinse both with water. See Figure L.

Allow to air dry.

Figure K. Removing the plunger from the barrel

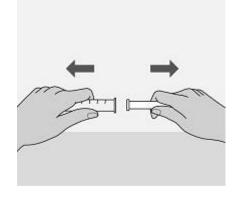
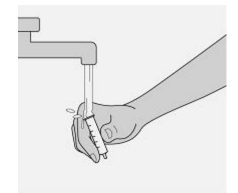


Figure L. Rinsing the plunger and barrel

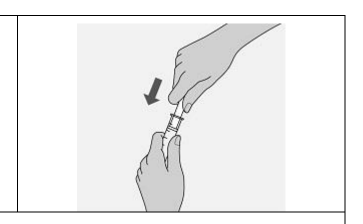


### Step 12. Put the oral syringe back together.

When the barrel and plunger are dry, put the oral syringe back together by inserting the plunger into the barrel. **See Figure M**. Store the oral syringe with the SELZENTRY oral solution.

Do not throw away the oral syringe.

Figure M. Putting the oral syringe back together



#### **How should I store SELZENTRY?**

Store SELZENTRY oral solution at room temperature between 68°F to 77°F (20°C to 25°C). Throw away any unused oral solution 60 days after first opening the bottle.

Keep SELZENTRY and all medicines out of the reach of children.

Manufactured for:



ViiV Healthcare

Research Triangle Park, NC 27709

SELZENTRY is a trademark owned by or licensed to the ViiV Healthcare group of companies.

©2018 ViiV Healthcare group of companies or its licensor.

SEL:2IFU

971

This Instructions for Use has been approved by the U.S. Food and Drug Administration.

Revised: 07/2018